FORM PTO-1390DEPARTMENT OF COMMERCE PATENT AND TRADEMARK OFFICE (REV 10-2000) ATTORNEY'S DOCKET NO. 11362.0030.PCUS00 (INNS:030) U.S. APPLICATION NO. (If known, see TRANSMITTAL LETTER TO THE UNITED STATES 37 CFR 1.5) DESIGNATED/ELECTED OFFICE (DO/EO/US) 09/720435 **CONCERNING A FILING UNDER 35 U.S.C. 371** PRIORITY DATE CLAIMED INTERNATIONAL APPLICATION NO. INTERNATIONAL FILING DATE 24 June 1998 22 June 1999 PCT/EP99/04317 TITLE OF INVENTION METHOD FOR DETECTION OF DRUG-SELECTED MUTATIONS IN THE HIV PROTEASE GENE APPLICANT(S) FOR DO/EO/US

App	ican	t herewith submits to the United States Designated/Elected Office (DO/EO/US) the following items and other information:
1.	$\bowtie$	This is a <b>FIRST</b> submission of items concerning a filing under 35 U.S.C. 371.
2.	$\overline{\Box}$	This is a <b>SECOND</b> or <b>SUBSEQUENT</b> submission of items concerning a filing under 35 U.S.C. 371.
3.		This express request to begin national examination procedures (35 U.S.C. 371(f)).
4.	$\overline{\boxtimes}$	The US has been elected by the expiration of 19 months from the priority date (PCT Article 31).
5.	$\overline{\boxtimes}$	A copy of the International Application as filed (35 U.S.C. 371(c)(2)).
		a.  is attached hereto (required only if not transmitted by the International Bureau).
		b.
		c. is not required, as the application was filed in the United States Receiving Office (RO/US).
6.		A translation of the International Application into English (35 U.S.C. 371(c)(2)).
7.	$\boxtimes$	Amendments to the claims of the International Application under PCT Article 19 (35 U.S.C. 371(c)(3)).
		a.  are attached hereto (required only if not transmitted by the International Bureau).
		b. have been communicated by the International Bureau.
		c. A have not been made; however, the time limit for making such amendments has NOT expired.
		d. have not been made and will not be made.
8.		A translation of the amendments to the claims under PCT Article 19 (35 U.S.C. 371(c)(3)).
9.		An oath or declaration of the inventor(s) (35 U.S.C. 371(c)(4)).
10.		An English translation of the annexes to the International Preliminary Examination Report under PCT Article 36 (35 U.S.C. 371(c)(5)).
Iten	ns 11	to 16 below concern document(s) or information included:
11.	$\boxtimes$	An Information Disclosure Statement under 37 CFR 1.97 and 1.98.
12.		An assignment document for recording. A separate cover sheet in compliance with 37 CFR 3.28 and 3.31 is included.
13.		A FIRST preliminary amendment. A SECOND or SUBSEQUENT preliminary amendment.
14.	$\boxtimes$	A substitute specification: Pages 8 and 33; and Figure 3 (See Rectification Notices dated January 11, 2000)
15.		A change of power of attorney and/or address letter.
16.	$\boxtimes$	Other items or information:
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#### 1. FIELD OF THE INVENTION

The present invention relates to the field of HIV diagnosis. More particularly, the present invention relates to the field of diagnosing the susceptibility of an HIV sample to antiviral drugs used to treat HIV infection.

The present invention relates to a method for the rapid and reliable detection of drug-selected mutations in the HIV protease gene allowing the simultaneous characterization of a range of codons involved in drug resistance using specific sets of probes optimized to function together in a reverse-hybridization assay.

#### 2. BACKGROUND OF THE INVENTION

The human immunodeficiency virus (HIV) is the ethiological agent for the acquired immunodeficiency syndrome (AIDS). HIV, like other retroviruses, encodes an aspartic protease that mediates the maturation of the newly produced viral particle by cleaving viral polypeptides into their functional forms (Hunter et al). The HIV protease is a dimeric molecule consisting of two identical subunits each contributing a catalytic aspartic residue (Navia et al, Whodawer et al, Meek et al). Inhibition of this enzyme gives rise to noninfectious viral particles that cannot establish new cycles of viral replication (Kohl et al, Peng et al).

Attempts to develop inhibitors of HIV-1 protease were initially based on designing peptide compounds that mimicked the natural substrate. The availability of the 3-dimensional structure of the enzyme have more recently allowed the rational design of protease inhibitors (PI) using computer modeling (Huff et al, Whodawer et al). A number of second generation PI that are partially peptidic or entirely nonpeptidic have proven to exhibit particularly potent antiviral effects in cell culture. Combinations of various protease inhibitors with nucleoside and non-nucleoside RT inhibitors have also been studied extensively in vitro. In every instance, the combinations have been at least additive and usually synergistic.

In spite of the antiviral potency of many recently developed HIV-1 PI, the emergence of virus variants with decreased sensitivity to these compounds has been described both in cell culture and in treated patients thereby escaping the inhibitory effect of the antiviral (Condra *et al.*). Emergence of

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resistant variants depends on the selective pressure applied to the viral population. In the case of a relatively ineffective drug, selective pressure is low because replication of both wild-type virus and any variants can continue. If a more effective drug suppresses replication of virus except for a resistant variant, then that variant will be selected. Virus variants that arise from selection by PI carry several distinct mutations in the protease coding sequence that appear to emerge sequentially. A number of these cluster near the active site of the enzyme while others are found at distant sites. This suggests conformational adaptation to primary changes in the active site and in this respect certain mutations that increase resistance to PI also decrease protease activity and virus replication.

Amongst the PI, the antiviral activity of the PI ritonavir (A-75925; ABT-538), nelfinavir (AG-1343), indinavir (MK-639; L735; L524) and saquinavir (Ro 31-8959) have been approved by the Food and Drug Administration and are currently under evaluation in clinical trials involving HIV-infected patients. The VX-487 (141W94) antiviral compound is not yet approved. The most important mutations selected for the above compounds and leading to gradually increasing resistance are found at amino acid (aa) positions 30 (D to N), 46 (M to I), 48 (G to V), 50 (I to V), 54 (I to A, I to V), 82 (V to A, or F, or I, or T), 84 (I to V) and 90 (L to M). Other mutations associated with drug resistance to the mentioned compounds have been described (Schinazi et al). Saquinavir-resistant variants, which usually carry mutations at amino acid positions 90 and/or 48, emerge in approximately 45% of patients after 1 year of monotherapy. Resistance appears to develop less frequently with higher doses of saquinavir. Resistance to indinavir and ritonavir requires multiple mutations; usually at greater than 3 and up to 11 sites, with more amino acid substitutions conferring higher levels of resistance. Resistant isolates usually carry mutations at codons 82, 84, or 90. In the case of ritonavir, the mutation at codon 82 appears first in most patients. Although mutant virions resistant to saquinavir are not cross-resistant to indinavir or ritonavir, isolates resistant to indinavir are generally ritonavir resistant and visa versa. Resistance to either indinavir or ritonavir usually results in cross-resistance to saquinavir. Approximately one third of indinavir resistant isolates are cross-resistant to nelfinavir as well.

The regime for an efficient antiviral treatment is currently not clear at all. Patterns of reduced susceptibility to HIV protease inhibitors have been investigated *in vitro* by cultivating virus in the presence of PI. These data, however, do not completely predict the pattern of amino-acid changes actually seen in patients receiving PI. Knowledge of the resistance and cross-resistance patterns should facilitate selection of optimal drug combinations and selection of sequences with non-overlapping resistance patterns. This would delay the emergence of cross-resistant viral strains and prolong the duration of effective antiretroviral activity in patients. Therefore, there is need for methods and systems that detect these mutational events in order to give a better insight into the mechanisms of HIV resistance. Further, there is need for methods and systems which can provide data important for the antiviral therapy to follow in a more time-efficient and economical manner compared to the conventional cell-culture selection techniques.

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#### 3. AIMS OF THE INVENTION

It is an aim of the present invention to develop a rapid and reliable detection method for determination of the antiviral drug resistance of viruses, which contain protease genes such as HIV retroviruses present in a biological sample.

More particularly it is an aim of the present invention to provide a genotyping assay allowing the detection of the different HIV protease gene wild type and mutation codons involved in the antiviral resistance in one single experiment.

It is also an aim of the present invention to provide an HIV protease genotyping assay or method which allows to infer the nucleotide sequence at codons of interest and/or the amino acids at the codons of interest and/or the antiviral drug selected spectrum, and possibly also infer the HIV type or subtype isolate involved.

Even more particularly it is an aim of the present invention to provide a genotyping assay allowing the detection of the different HIV protease gene polymorphisms representing wild-type and mutation codons in one single experimental setup.

It is another aim of the present invention to select particular probes able to discriminate wild-type HIV protease sequences from mutated or polymorphic HIV protease sequences conferring resistance to one or more antiviral drugs, such as ritonavir (A-75925; ABT-538), nelfinavir (AG-1343), indinavir (MK-639; L735; L524), saquinavir (Ro 31-8959) and VX-478 (141W94) or others (Shinazi *et al*).

It is more particularly an aim of the present invention to select particular probes able to discriminate wild-type HIV protease sequences from mutated or polymorphic HIV protease sequences conferring resistance to ritonavir (A-75925; ABT-538).

It is more particularly an aim of the present invention to select particular probes able to discriminate wild-type HIV protease sequences from mutated HIV protease sequences conferring resistance to nelfinavir (AG-1343).

It is more particularly an aim of the present invention to select particular probes able to discriminate wild-type HIV protease sequences from mutated HIV protease sequences conferring resistance to indinavir (MK-639; L735; L524).

It is more particularly an aim of the present invention to select particular probes able to discriminate wild-type HIV protease sequences from mutated HIV protease sequences conferring resistance to saquinavir (Ro 31-8959).

It is more particularly an aim of the present invention to select particular probes able to discriminate wild-type HIV protease sequences from mutated HIV protease sequences conferring resistance to VX-478 (141W94).

It is also an aim of the present invention to select particular probes able to determine and/or infer

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cross-resistance to HIV protease inhibitors.

It is more particularly an aim of the present invention to select particular probes able to discriminate wild-type HIV protease from mutated HIV protease sequences involving at least one of amino acid positions 30 (D to N), 46 (M to I), 48 (G to V), 50 (I to V), 54 (I to A or V), 82 (V to A or F or I or T), 84(I to V) and 90 (L to M) of the viral protease gene.

It is particularly an aim of the present invention to select a particular set of probes, able to discriminate wild-type HIV protease sequences from mutated HIV protease sequences conferring resistance to any of the antiviral drugs defined above with this particular set of probes being used in a reverse hybridization assay.

It is moreover an aim of the present invention to combine a set of selected probes able to discriminate wild-type HIV protease sequences from mutated HIV protease sequences conferring resistance to antiviral drugs with another set of selected probes able to identify the HIV isolate, type or subtype present in the biological sample, whereby all probes can be used under the same hybridization and wash-conditions.

It is also an aim of the present invention to select primers enabling the amplification of the gene fragment(s) determining the antiviral drug resistance trait of interest.

It is also an aim of the present invention to select particular probes able to identify mutated HIV protease sequences resulting in cross-resistance to antiviral drugs.

The present invention also aims at diagnostic kits comprising said probes useful for developing such a genotyping assay.

The present invention also aims at diagnostic kits comprising said primers useful for developing such a genotyping assay.

### 4. DETAILED DESCRIPTION OF THE INVENTION.

All the aims of the present invention have been met by the following specific embodiments.

According to one embodiment, the present invention relates to a method for determining the susceptibility to antiviral drugs of HIV viruses in a biological sample, with said method comprising:

- a) if need be, releasing, isolating or concentrating the polynucleic acids present in the sample;
- b) if need be amplifying the relevant part of the protease gene of HIV with at least one suitable primer pair;
- c) hybridizing the polynucleic acids of step a) or b) with at least one of the following probes:
  - probes specifically hybridizing to a target sequence comprising codon 30;
- probes specifically hybridizing to a target sequence comprising codon 46 and/or 48; probes specifically hybridizing to a target sequence comprising codon 50;

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probes specifically hybridizing to a target sequence comprising codon 54; probes specifically hybridizing to a target sequence comprising codon 82 and/or 84; probes specifically hybridizing to a target sequence comprising codon 90; or the complement of said probes,

further characterized in that said probes specifically hybridize to any of the target sequences presented in figure 1, or to the complement of said target sequences;

d) inferring from the result of step c) whether or not a mutation giving rise to drug resistance is present in any of said target sequences.

The numbering of HIV-1 protease gene encoded amino acids is as generally accepted in literature.

Mutations that give rise to an amino acid change at position 48 or 90 are known to confer resistance to saquinavir (Erlebe et al; Tisdale et al). An amino acid change at codon 46 or 54 or 82 or 84 results in ritonavir and indinavir resistance (Kempf et al; Emini et al; Condra et al). Amino acid changes at positions 30 and 46 confer resistance to nelfinavir (Patick et al) and amino acid changes at position 50 confers resistance to VX-487 (Rao et al). Therefore, the method described above allows to determine whether a HIV strain is susceptible or resistant to any of the drugs mentioned above. This method can be used, for instance, to screen for mutations conferring resistance to any of the mentioned drugs before initiating therapy. This method may also be used to screen for mutations that may arise during the course of therapy (i.e. monitoring of drug therapy). It is obvious that this method may also be used to determine resistance to drugs other than the above-mentioned drugs, provided that resistance to these other drugs is linked to mutations that can be detected by use of this method. This method may also be used for the specific detection of polymorphic nucleotides. It is to be understood that the said probes may only partly overlap with the targets sequences of figure 1, table 2 and table 3, as long as they allow for specific detection of the relevant polymorphic nucleotides as indicated above. The sequences of figure 1, table 2 and table 3 were derived from polynucleic acid fragments comprising the protease gene. These fragments were obtained by PCR amplification and were inserted into a cloning vector and sequence analyzed as described in example 1. It is to be noted that some polynucleic acid fragments comprised polymorphic nucleotides in their sequences, which have not been previously disclosed. These novel polymorphic nucleotide sequences are represented in table 4 below.

TABLE 4: Polymorphic nucleotide sequences.

	51	52	53	54	55	56	57	58	code	n r	osi	tion
	gga	ggt	ttt	atc	aaa	gta	aga	cag	cons	ens	us	sequence
	GGA	GGT	TTT	ATC	AAA	GTC	AGA	CAA	SEQ	ID	NO	478
35	GGA	GGT	TTC	ATT	AAG	GTA	AAA	CAG	SEQ	ID	ИО	479
	GGA	GGT	TTT	ATT	AAG	GTA	AGA	CAG	SEQ	ID	ио	480

GGA GGT TTT ATT AAA GTA AGA CAA

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SEQ ID NO 481

	GGA	GGC	TTT	ATC	AAA	GTA	AGA	CAA		SEQ	ID	NO	482
	GGA	GGT	TTT	ATC	AAA	GTC	AGA	CAA		SEQ	ID	ИО	483
5	78	79	80	81	82	83	84	85		code	n p	osi	tion
	gga	cct	aca	cct	gtc	aac	ata	att	gg	cons	ens	sus	sequence
	GGA	CCT	ACA	CCG	GTC	AAC	ATA	ATT	GG	SEQ	ID	ИО	484
	GGA	CCT	ACA	CCT	GCC	AAT	ATA	ATT	GG	SEQ	ID	ио	485
	GGA	CCT	ACG	CCC	TTC	AAC	ATA	ATT	GG	SEQ	ID	NO	486
10	GGA	CCG	ACA	CCT	GTC	ACC	ATA	ATT	GG	SEQ	ID	ио	487
	GGA	CCT	ATA	CCT	GTC	AAC	ATA	ATT	GG	SEQ	ID	ио	488

	87	88	89	90	91	92	93	94	codon position
a	aga	aat	ctg	ttg	act	cag	att	ggc	consensus sequence
A	AAA	TAA	CTG	ATG	ACT	CAG	ATT	GGC	SEQ ID NO 489
Α	AGA	ACT	CTG	TTG	ACT	CAG	CTT	GGA	SEQ ID NO 490
A	AGA	AAT	ATG	ATG	ACC	CAG	CTT	GGC	SEQ ID NO 491
A	AGA	AAT	ATA	ATG	ACT	CAG	CTT	GGA	SEQ ID NO 492
Α	AGA	AAT	CTG	CTG	ACT	CAG	ATT	GGG	SEQ ID NO 493
A	AGA	AAT	CTG	TTG	ACA	CAG	CTT	GGC	SEQ ID NO 494
A	AGA	AAT	ATG	TTG	ACT	CAG	CTT	GGT	SEQ ID NO 495
A	AGA	AAT	TTG	TTG	ACT	CAG	ATT	GGG	SEQ ID NO 496
A	AGA	AAT	ATG	TTG	ACT	CAG	CTT	GGT	SEQ ID NO 497
Α	AGA	AAT	ATG	TTG	ACT	CAG	CTT	GGA	SEQ ID NO 498
Α	AGA	AAT	CTG	TTG	ACT	CAG	CTT	GGA	SEQ ID NO 499
Α	. AGA	AAC	CTG	TTG	ACT	CAA	CTT	GGT	SEQ ID NO 500

The present invention thus also relates to these novel sequences, or a fragment thereof, wherein said fragment consists of at least 10, preferably 15, even more preferably 20 contiguous nucleotides and contains at least one polymorphic nucleotide. It is furthermore to be understood that these new

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polymorphic nucleotides may also be expected to arise in another sequence context than in the mentioned sequences. For instance a G at the third position of codon 55 is shown in SEQ ID N° 478 in combination with a T at the third position of codon 54, but a G at the third position of codon 55 may also be expected to occur in the context of a wild type sequence. It is also to be understood that the above mentioned specifications apply to the complement of the said target sequences as well. This applies also to Figure 1.

According to a preferred embodiment the present invention relates to a method as indicated above, further characterized in that said probes are capable of simultaneously hybridizing to their respective target regions under appropriate hybridization and wash conditions allowing the detection of the hybrids formed.

According to a preferred embodiment, step c is performed using a set of at least 2, preferably at least 3, more preferably at least 4 and most preferably at least 5 probes meticulously designed as such that they show the desired hybridization results. In general this method may be used for any purpose that relies on the presence or absence of mutations that can be detected by this method, e.g. for genotyping. The probes of table 1 have been optimized to give specific hybridization results when used in a LiPA assay (see below), as described in examples 2 and 3. These probes have thus also been optimized to simultaneously hybridize to their respective target regions under the same hybridization and wash conditions allowing the detection of hybrids. The sets of probes for each of the codons 30, 46/48, 50, 54 and 82/84 have been tested experimentally as described in examples 2 and 3. The reactivity of the sets shown in table 1 with 856 serum samples from various geographic origins was evaluated. It was found that the sets of probes for codons 30, 46/48, 50, 54 and 82/84 reacted with 98.9%, 99.6%, 98.5%, 99.2%, 95.4% and 97.2% of the test samples, respectively. The present invention thus also relates to the sets of probes for codons 30, 46/48, 50, 54, 82/84 and 90, shown in table 1 and table 7.

According to another even more preferred embodiment, the present invention relates to a method as defined above, further characterized in that:

step b) comprises amplifying a fragment of the protease gene with at least one 5'-primer specifically hybridizing to a target sequence located between nucleotide position 210 and nucleotide position 260 (codon 87), more preferably between nucleotide position 220 and nucleotide position 260 (codon 87), more preferably between nucleotide position 230 and nucleotide position 260 (codon 87), even more preferably at nucleotide position 241 to nucleotide position 260 (codon 87) in combination with at least one suitable 3'-primer, and

step c) comprises hybridizing the polynucleic acids of step a) or b) with at least one of the probes specifically hybridizing to a target sequence comprising codon 90.

According to another even more preferred embodiment, the present invention relates to a method as defined above, further characterized in that:

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step b) comprises amplifying a fragment of the protease gene with at least one 3'-primer specifically hybridizing to a target sequence located between nucleotide position 253 (codon 85) and nucleotide positions 300, more preferably between nucleotide position 253 (codon 85) and nucleotide positions 290, more preferably between nucleotide position 253 (codon 85) and nucleotide positions 280, even more preferably at nucleotide position 253 (codon 85) to nucleotide position 273 (codon 91), in combination with at least one suitable 5'-primer, and step c) comprises hybridizing the polynucleic acids of step a) or b) with at least one of the probes specifically hybridizing to a target sequence comprising any of codons 30, 46, 48, 50, 52, 54, 82 and 84.

It has been found, unexpectedly, that an amplified nucleic acid fragment comprising all of the above-mentioned codons, does not hybridize optimally to probes comprising codon 82, 84 or 90. On the other hand, a shorter fragment, for instance the fragment which is amplified by use of the primers Prot41bio and Prot6bio with respectively seq id no 5 and seq id no 4, hybridizes better to probes comprising codon 90. Better hybridization is also obtained when the fragment is amplified with primer Prot41bio in combination with primers Prot6abio, Prot6bbio, Prot6cbio and Prot6dbio The present invention thus also relates to a method as defined above, further characterized in that the 5'-primer is seq id no 5 and at least one 3' primer is chosen from seq id no 4, seq id no506, seq id no 507, seq id no 508, and seq id no 509. Likewise, another shorter fragment, for instance the fragment which is amplified by use of the primers Prot2bio and Prot31bio with respectively seq id no 3 and seq id no 6, was found to hybridize better to probes comprising codon 82 and/or 84. Hence the present invention also relates to a method as defined above, further characterized in that the 5'-primer is seq id no 5 and at least one 3'-primer is chosen from seq id no 4, seq id no 506, seq id no 507, seq id no 508, and seq id no 509.

New sets of amplification primers as mentioned in example 1 were selected. The present invention thus also relates to primers: prot 16 (SEQ ID NO 501), prot 5 (SEQ ID NO 5), prot2a bio (SEQ ID NO 503), prot2b bio (SEQ ID NO 504), prot31 bio (SEQ ID NO 6), prot41-bio (SEQ ID NO 505), prot6a (SEQ ID NO 506), prot6b (SEQ ID NO 507), prot6c (SEQ ID NO 508) and prot6d (SEQ ID NO 509). A number of these primers are chemically modified (biotinylated), others are not. The present invention relates to any of the primers mentioned, primers containing unmodified nucleotides, or primers containing modified nucleotides.

Different techniques can be applied to perform the sequence-specific hybridization methods of the present invention. These techniques may comprise immobilizing the amplified HIV polynucleic acids on a solid support and performing hybridization with labeled oligonucleotide probes. HIV polynucleic acids may also be immobilized on a solid support without prior amplification and subjected to hybridization. Alternatively, the probes may be immobilized on a solid support and hybridization may be performed with labeled HIV polynucleic acids, preferably after amplification. This technique is called reverse hybridization. A convenient reverse hybridization technique is the line probe assay (LiPA). This

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assay uses oligonucleotide probes immobilized as parallel lines on a solid support strip (Stuyver et al., 1993). It is to be understood that any other technique based on the above-mentioned methods is also covered by the present invention.

According to another preferred embodiment, the present invention relates to any of the probes mentioned above and/or to any of the primers mentioned above, with said primers and probes being designed for use in a method for determining the susceptibility to antiviral drugs of HIV viruses in a sample. According to an even more preferred embodiment, the present invention relates to the probes with seq id no 7 to seq id no 477 and seq id no510 to seq id no 519, more preferably to the seq id no mentioned in Table 1 and Table 7, and to the primers with seq id no 3, 4, 5 and 6, 501, 502, 503, 504, 505, 506, 507, 508 and 509. The skilled man will recognize that addition or deletion of one or more nucleotides at their extremities may adapt the said probes and primers. Such adaptations may be required if the conditions of amplification or hybridization are changed, or if the amplified material is RNA instead of DNA, as is the case in the NASBA system.

According to another preferred embodiment, the present invention relates to a diagnostic kit enabling a method for determining the susceptibility to antiviral drugs of HIV viruses in a biological sample, with said kit comprising:

- a) when appropriate, a means for releasing, isolating or concentrating the polynucleic acids present in said sample;
- b) when appropriate, at least one of the primers of any of claims 4 to 6;
- c) at least one of the probes of any of claims 1 to 3, possibly fixed to a solid support;
  - d) a hybridization buffer, or components necessary for producing said buffer;
  - e) a wash solution, or components necessary for producing said solution;
  - f) when appropriate, a means for detecting the hybrids resulting from the preceding hybridization;
  - h) when appropriate, a means for attaching said probe to a solid support.

#### **DEFINITIONS**

The following definitions serve to illustrate the terms and expressions used in the present invention.

The term "antiviral drugs" refers particularly to any antiviral protease inhibitor. Examples of such antiviral drugs and the mutation they may cause in the HIV protease gene are disclosed in Schinazi et al., 1997. The contents of the latter two documents particularly are to be considered as forming part of the present invention. The most important antiviral drugs focussed at in the present invention are disclosed in Tables 1 to 2.

The target material in the samples to be analyzed may either be DNA or RNA, e.g.: genomic DNA, messenger RNA, viral RNA or amplified versions thereof. These molecules are also termed polynucleic acids.

It is possible to use genomic DNA or RNA molecules from HIV samples in the methods according to the present invention.

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Well-known extraction and purification procedures are available for the isolation of RNA or DNA from a sample (fi. in Maniatis et al., Molecular Cloning: A Laboratory Manual, 2nd Edition, Cold Spring Harbour Laboratory Press (1989)).

The term "probe" refers to single stranded sequence-specific oligonucleotides, which have a sequence, which is complementary to the target sequence to be detected.

The term "target sequence" as referred to in the present invention describes the wild type nucleotide sequence, or the sequence comprising one or more polymorphic nucleotides of the protease gene to be specifically detected by a probe according to the present invention. This nucleotide sequence may encompass one or several nucleotide changes. Target sequences may refer to single nucleotide positions, codon positions, nucleotides encoding amino acids or to sequences spanning any of the foregoing nucleotide positions. In the present invention said target sequence often includes one or two variable nucleotide positions.

The term "polymorphic nucleotide" indicates a nucleotide in the protease gene of a particular HIV virus that is different from the nucleotide at the corresponding position in at least one other HIV virus. The polymorphic nucleotide may or may not give rise to resistance to an antiviral drug. It is to be understood that the complement of said target sequence is also a suitable target sequence in some cases. The target sequences as defined in the present invention provide sequences which should be complementary to the central part of the probe which is designed to hybridize specifically to said target region.

The term "complementary" as used herein means that the sequence of the single stranded probe is exactly the (inverse) complement of the sequence of the single-stranded target, with the target being defined as the sequence where the mutation to be detected is located.

"Specific hybridization" of a probe to a target sequence of the HIV polynucleic acids means that said probe forms a duplex with part of this region or with the entire region under the experimental conditions used, and that under those conditions said probe does not form a duplex with other regions of the polynucleic acids present in the sample to be analyzed.

Since the current application requires the detection of single basepair mismatches, very stringent conditions for hybridization are required, allowing in principle only hybridization of exactly complementary sequences. However, variations are possible in the length of the probes (see below), and it should be noted that, since the central part of the probe is essential for its hybridization characteristics, possible deviations of the probe sequence versus the target sequence may be allowable towards head and tail of the probe, when longer probe sequences are used. These variations, which may be conceived from the common knowledge in the art, should however always be evaluated experimentally, in order to check if they result in equivalent hybridization characteristics than the exactly complementary probes.

Preferably, the probes of the invention are about 5 to 50 nucleotides long, more preferably from about 10 to 25 nucleotides. Particularly preferred lengths of probes include 10, 11, 12, 13, 14, 15, 16, 17,

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18, 19, 20, 21, 22, 23, 24 or 25 nucleotides. The nucleotides as used in the present invention may be ribonucleotides, deoxyribonucleotides and modified nucleotides such as inosine or nucleotides containing modified groups, which do not essentially alter their hybridization characteristics.

Probe sequences are represented throughout the specification as single stranded DNA oligonucleotides from the 5' to the 3' end. It is obvious to the man skilled in the art that any of the below-specified probes can be used as such, or in their complementary form, or in their RNA form (wherein U replaces T).

The probes according to the invention can be prepared by cloning of recombinant plasmids containing inserts including the corresponding nucleotide sequences, if need be by cleaving the latter out from the cloned plasmids upon using the adequate nucleases and recovering them, e.g. by fractionation according to molecular weight. The probes according to the present invention can also be synthesized chemically, for instance by the conventional phospho-triester method.

The term "solid support" can refer to any substrate to which an oligonucleotide probe can be coupled, provided that it retains its hybridization characteristics and provided that the background level of hybridization remains low. Usually the solid substrate will be a microtiter plate, a membrane (e.g. nylon or nitrocellulose) or a microsphere (bead) or a chip. Prior to application to the membrane or fixation it may be convenient to modify the nucleic acid probe in order to facilitate fixation or improve the hybridization efficiency. Such modifications may encompass homopolymer tailing, coupling with different reactive groups such as aliphatic groups, NH<sub>2</sub> groups, SH groups, carboxylic groups, or coupling with biotin, haptens or proteins.

The term "labeled" refers to the use of labeled nucleic acids. Labeling may be carried out by the use of labeled nucleotides incorporated during the polymerase step of the amplification such as illustrated by Saiki et al. (1988) or Bej et al. (1990) or labeled primers, or by any other method known to the person skilled in the art. The nature of the label may be isotopic (<sup>32</sup>P, <sup>35</sup>S, etc.) or non-isotopic (biotin, digoxigenin, etc.).

The term "primer" refers to a single stranded oligonucleotide sequence capable of acting as a point of initiation for synthesis of a primer extension product, which is complementary to the nucleic acid strand to be copied. The length and the sequence of the primer must be such that they allow to prime the synthesis of the extension products. Preferably the primer is about 5-50 nucleotides long. Specific length and sequence will depend on the complexity of the required DNA or RNA targets, as well as on the conditions of primer use such as temperature and ionic strength.

The term "primer pair" refers to a set of primers comprising at least one 5' primer and one 3' primer. The primer pair may consist of more than two primers, the complexity of the number of primers will depend on the hybridization conditions, variability of the sequences in the regions to be amplified and the target sequences to be detected.

The fact that amplification primers do not have to match exactly with the corresponding template

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sequence to warrant proper amplification is amply documented in the literature (Kwok et al., 1990).

The amplification method used can be either polymerase chain reaction (PCR; Saiki et al., 1988), ligase chain reaction (LCR; Landgren et al., 1988; Wu & Wallace, 1989; Barany, 1991), nucleic acid sequence-based amplification (NASBA; Guatelli et al., 1990; Compton, 1991), transcription-based amplification system (TAS; Kwoh et al., 1989), strand displacement amplification (SDA; Duck, 1990) or amplification by means of Qß replicase (Lomeli et al., 1989) or any other suitable method to amplify nucleic acid molecules known in the art.

The oligonucleotides used as primers or probes may also comprise nucleotide analogues such as phosphorothiates (Matsukura et al., 1987), alkylphosphorothiates (Miller et al., 1979) or peptide nucleic acids (Nielsen et al., 1991; Nielsen et al., 1993) or may contain intercalating agents (Asseline et al., 1984).

As most other variations or modifications introduced into the original DNA sequences of the invention these variations will necessitate adaptations with respect to the conditions under which the oligonucleotide should be used to obtain the required specificity and sensitivity. However the eventual results of hybridization will be essentially the same as those obtained with the unmodified oligonucleotides.

The introduction of these modifications may be advantageous in order to positively influence characteristics such as hybridization kinetics, reversibility of the hybrid-formation, biological stability of the oligonucleotide molecules, etc.

The "sample" may be any biological material taken either directly from the infected human being (or animal), or after culturing (enrichment). Biological material may be e.g. expectorations of any kind, broncheolavages, blood, skin tissue, biopsies, sperm, lymphocyte blood culture material, colonies, liquid cultures, fecal samples, urine etc.

The sets of probes of the present invention will include at least 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30 or more probes. Said probes may be applied in two or more distinct and known positions on a solid substrate. Often it is preferable to apply two or more probes together in one and the same position of said solid support.

For designing probes with desired characteristics, the following useful guidelines known to the person skilled in the art can be applied.

Because the extent and specificity of hybridization reactions such as those described herein are affected by a number of factors, manipulation of one or more of those factors will determine the exact sensitivity and specificity of a particular probe, whether perfectly complementary to its target or not. The importance and effect of various assay conditions, explained further herein, are known to those skilled in the art.

The stability of the [probe: target] nucleic acid hybrid should be chosen to be compatible with the assay conditions. This may be accomplished by avoiding long AT-rich sequences, by terminating the

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hybrids with G:C base pairs, and by designing the probe with an appropriate Tm. The beginning and end points of the probe should be chosen so that the length and %GC result in a Tm about 2-10°C higher than the temperature at which the final assay will be performed. The base composition of the probe is significant because G-C base pairs exhibit greater thermal stability as compared to A-T base pairs due to additional hydrogen bonding. Thus, hybridization involving complementary nucleic acids of higher G-C content will be stable at higher temperatures.

Conditions such as ionic strength and incubation temperature under which a probe will be used should also be taken into account when designing a probe. It is known that hybridization will increase as the ionic strength of the reaction mixture increases, and that the thermal stability of the hybrids will increase with increasing ionic strength. On the other hand, chemical reagents, such as formamide, urea, DMSO and alcohols, which disrupt hydrogen bonds, will increase the stringency of hybridization. Destabilization of the hydrogen bonds by such reagents can greatly reduce the Tm. In general, optimal hybridization for synthetic oligonucleotide probes of about 10-50 bases in length occurs approximately 5°C below the melting temperature for a given duplex. Incubation at temperatures below the optimum may allow mismatched base sequences to hybridize and can therefore result in reduced specificity.

It is desirable to have probes, which hybridize only under conditions of high stringency. Under high stringency conditions only highly complementary nucleic acid hybrids will form; hybrids without a sufficient degree of complementarity will not form. Accordingly, the stringency of the assay conditions determines the amount of complementarity needed between two nucleic acid strands forming a hybrid. The degree of stringency is chosen such as to maximize the difference in stability between the hybrid formed with the target and the nontarget nucleic acid. In the present case, single base pair changes need to be detected, which requires conditions of very high stringency.

The length of the target nucleic acid sequence and, accordingly, the length of the probe sequence can also be important. In some cases, there may be several sequences from a particular region, varying in location and length, which will yield probes with the desired hybridization characteristics. In other cases, one sequence may be significantly better than another that differs merely by a single base. While it is possible for nucleic acids that are not perfectly complementary to hybridize, the longest stretch of perfectly complementary base sequence will normally primarily determine hybrid stability. While oligonucleotide probes of different lengths and base composition may be used, preferred oligonucleotide probes of this invention are between about 5 to 50 (more particularly 10-25) bases in length and have a sufficient stretch in the sequence which is perfectly complementary to the target nucleic acid sequence.

Regions in the target DNA or RNA, which are known to form strong internal structures inhibitory to hybridization, are less preferred. Likewise, probes with extensive self-complementarity should be avoided. As explained above, hybridization is the association of two single strands of complementary nucleic acids to form a hydrogen bonded double strand. It is implicit that if one of the two strands is wholly or partially involved in a hybrid that it will be less able to participate in formation

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of a new hybrid. There can be intramolecular and intermolecular hybrids formed within the molecules of one type of probe if there is sufficient self complementarity. Such structures can be avoided through careful probe design. By designing a probe so that a substantial portion of the sequence of interest is single stranded, the rate and extent of hybridization may be greatly increased. Computer programs are available to search for this type of interaction. However, in certain instances, it may not be possible to avoid this type of interaction.

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Standard hybridization and wash conditions are disclosed in the Materials & Methods section of the Examples. Other conditions are for instance 3X SSC (Sodium Saline Citrate), 20% deionized FA (Formamide) at 50°C.

Other solutions (SSPE (Sodium saline phosphate EDTA), TMACI (Tetramethyl ammonium Chloride), etc.) and temperatures can also be used provided that the specificity and sensitivity of the probes is maintained. If need be, slight modifications of the probes in length or in sequence have to be carried out to maintain the specificity and sensitivity required under the given circumstances.

Primers may be labeled with a label of choice (e.g. biotin). Different primer-based target amplification systems may be used, and preferably PCR-amplification, as set out in the examples. Single-round or nested PCR may be used.

The term "hybridization buffer" means a buffer enabling a hybridization reaction to occur between the probes and the polynucleic acids present in the sample, or the amplified products, under the appropriate stringency conditions.

The term "wash solution" means a solution enabling washing of the hybrids formed under the appropriate stringency conditions.

The following examples only serve to illustrate the present invention. These examples are in no way intended to limit the scope of the present invention.

#### 25 FIGURE AND TABLE LEGENDS

Figure 1: Natural and drug selected variability in the vicinity of codons 30, 46, 48, 50, 54, 82, 84, and 90 of the HIV-1 protease gene. The most frequently observed wild-type sequence is shown in the top line. Naturally occurring variations are indicated below and occur independently from each other. Drug-selected variants are indicated in bold

Figure 2 A: Reactivities of the selected probes for codon 30 immobilized on LiPA strips with reference material. The information in the boxed surface is not relevant for the discussion of probes for condon 30. The position of each selected probe on the membrane strip is shown at the left of each panel. The sequence of the relevant part of the selected probes is shown at the left and is given in Table 1. Each strip is incubated with a biotinylated PCR fragment from the reference panel. The reference panel accession

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numbers are indicated in <u>Table 1</u>. For several probes multiple reference panel possibilities are available, but only one relevant accession number given each time. \*: False positive reactivities. At the bottom the strips, the amino acids at the relevant codon, as derived from the probe reactivity, is indicated.

- Figure 2 B: Reactivities of the selected probes for codons 46 and 48 immobilized on LiPA strips with reference material. The information in the boxed surface is not relevant for the discussion of probes for condons 46 and 48 The position of each selected probe on the membrane strip is shown at the left of each panel. The sequence of the relevant part of the selected probes is given in Table 1. Each strip is incubated with a biotinylated PCR fragment from the reference panel. The reference panel accession numbers are indicated in <u>Table 1</u>. For several probes multiple reference panel possibilities are available, but only one relevant accession number given each time. \*: False positive reactivities. On top of the strips, the amino acids at the relevant codon, as derived from the probe reactivity, is indicated.
  - Figure 2 C: Reactivities of the selected probes for codon 50 immobilized on LiPA strips with reference material. The information in the boxed surface is not relevant for the discussion of probes for condon 50. The position of each selected probe on the membrane strip is shown at the left of each panel. The sequence of the relevant part of the selected probes is given in Table 1. Each strip is incubated with a biotinylated PCR fragment from the reference panel. The reference panel accession numbers are indicated in <u>Table 1</u>. For several probes multiple reference panel possibilities are available, but only one relevant accession number given each time. \*: False positive reactivities. At the bottom of the strips, the amino acids at the relevant codon, as derived from the probe reactivity, is indicated.
- Figure 2 D: Reactivities of the selected probes for codon 54 immobilized on LiPA strips with reference material. The information in the boxed surface is not relevant for the discussion of probes for condon 54.

  The position of each selected probe on the membrane strip is shown at the left of each panel. The sequence of the relevant part of the selected probes is given in Table 1. Each strip is incubated with a biotinylated PCR fragment from the reference panel. The reference panel accession numbers are indicated in Table 1. For several probes multiple reference panel possibilities are available, but only one relevant accession number given each time. \*: False positive reactivities. At the bottom of the strips, the amino acids at the relevant codon, as derived from the probe reactivity, is indicated.
  - Figure 2 E.:Reactivities of the selected probes for codons 82 and 84 immobilized on LiPA strips with reference material. The information in the boxed surface is not relevant for the discussion of probes for condons 82 and 84. The position of each selected probe on the membrane strip is shown at the left of each panel. The sequence of the relevant part of the selected probes is given in Table 1. Each strip is incubated with a biotinylated PCR fragment from the reference panel. The reference panel accession

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numbers are indicated in <u>Table 1</u>. For several probes multiple reference panel possibilities are available, but only one relevant accession number given each time. \*: False positive reactivities. At the bottom of the strips, the amino acids at the relevant codon, as derived from the probe reactivity, is indicated.

- Figure 2 F: Reactivities of the selected probes for codon 90 immobilized on LiPA strips with reference material. The information in the boxed surface is not relevant for the discussion of probes for condon 90.

  The position of each selected probe on the membrane strip is shown at the left of each panel. The sequence of the relevant part of the selected probes is given in Table 1. Each strip is incubated with a biotinylated PCR fragment from the reference panel. The reference panel accession numbers are indicated in Table 1. For several probes multiple reference panel possibilities are available, but only one relevant accession number given each time. \*: False positive reactivities. At the bottom of the strips, the amino acids at the relevant codon, as derived from the probe reactivity, is indicated.
  - Figure 3: Sequence and position of the HIV-1 protease amplification primers. To obtain the reactivity with probes selected to determine the susceptibility to antiviral drugs involving codons 30, 46, 48, 50, 54, 82, and 84, nested amplification primers prot2bio(5' primer) and Prot31bio (3' primer) were designed. To obtain the reactivity with probes selected to determine the susceptibility to antiviral drugs involving codon 90, nested amplification primers Prot41bio (5' primer) and Prot6bio (3' primer) were designed.
- Figure 4 A: Phylogenetic analysis on 312 protease sequences allowed to separate genotype B strains from non-B strains. Reactivities of the selected probes for codon 30 immobilized on LiPA strips with a biotinylated PCR fragment of genotype B strains and non-B strains is shown, the exact percentages are indicated in table 5. The probes are indicated at the bottom. The sequence of the relevant part of the probes is given in Table 1.

Figure 4 B: Phylogenetic analysis on 312 protease sequences allowed to separate genotype B strains from non-B strains. Reactivities of the selected probes for codons 46/48 immobilized on LiPA strips with a biotinylated PCR fragment of genotype B strains and non-B strains is shown, the exact percentages are indicated in table 5. The probes are indicated at the bottom. The sequence of the relevant part of the probes is given in Table 1.

Figure 4 C: Phylogenetic analysis on 312 protease sequences allowed to separate genotype B strains from non-B strains. Reactivities of the selected probes for codon 50 immobilized on LiPA strips with a biotinylated PCR fragment of genotype B strains and non-B strains is shown, the exact percentages are indicated in table 5. The probes are indicated at the bottom. The sequence of the relevant part of the probes is given in Table 1.

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Figure 4 D: Phylogenetic analysis on 312 protease sequences allowed to separate genotype B strains from non-B strains. Reactivities of the selected probes for codon 54 immobilized on LiPA strips with a biotinylated PCR fragment of genotype B strains and non-B strains is shown, the exact percentages are indicated in table 5. The probes are indicated at the bottom. The sequence of the relevant part of the probes is given in Table 1.

Figure 4 E: Phylogenetic analysis on 312 protease sequences allowed to separate genotype B strains from non-B strains. Reactivities of the selected probes for codons 82/84 immobilized on LiPA strips with a biotinylated PCR fragment of genotype B strains and non-B strains is shown, the exact percentages are indicated in table 5. The probes are indicated at the bottom. The sequence of the relevant part of the probes is given in Table 1.

Figure 4 F: Phylogenetic analysis on 312 protease sequences allowed to separate genotype B strains from non-B strains. Reactivities of the selected probes for codon 90 immobilized on LiPA strips with a biotinylated PCR fragment of genotype B strains and non-B strains is shown, the exact percentages are indicated in table 5. The probes are indicated at the bottom. The sequence of the relevant part of the probes is given in Table 1.

Figure 5 A: Geographical origin of 856 samples and reactivities with the different probes at codon position 30. The exact percentages are indicated in table 6. The probes are indicated at the bottom.

Figure 5 B: Geographical origin of 856 samples and reactivities with the different probes at codon positions 46/48. The exact percentages are indicated in table 6. The probes are indicated at the bottom.

Figure 5 C: Geographical origin of 856 samples and reactivities with the different probes at codon position 50. The exact percentages are indicated in table 6. The probes are indicated at the bottom.

Figure 5 D: Geographical origin of 856 samples and reactivities with the different probes at codon position 54. The exact percentages are indicated in table 6. The probes are indicated at the bottom.

Figure 5 E: Geographical origin of 856 samples and reactivities with the different probes at codon positions 82/84. The exact percentages are indicated in table 6. The probes are indicated at the bottom.

Figure 5 F: Geographical origin of 856 samples and reactivities with the different probes at codon position 90. The exact percentages are indicated in table 6. The probes are indicated at the bottom.

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Table 1: HIV-1 protease wild-type and drug-selected mutation probes with their corresponding sequences as applied on the HIV-1 protease LiPA strip. The most frequently observed wild-type sequence is shown at the top line. Probe names corresponding to the selected motifs are indicated in the left column, the relevant part of each probe applied on the strip is shown under the consensus sequence.

Table 2: Protease Inhibitors.

Table 3: HIV-1 protease wild-type and drug-selected mutation probes with their corresponding sequences as synthesized, immobilized and tested on LiPA strips. The most frequently observed wild-type sequence is shown at the top line. Probe names corresponding to the selected motifs are indicated in the left column, the relevant part of each probe applied on the strip is shown under the consensus sequence. The probes retained are indicated in table 1.

15 Table 4: Polymorphic nucleotide sequences.

Table 5: % Reactivities of the HIV-1 protease wild-type and drug-selected mutation probes applied on the HIV-1 protease LiPA strip with genotype B strains and non-B strains.

Table 6: % Reactivities of the HIV-1 protease wild-type and drug-selected mutation probes applied on the HIV-1 protease LiPA strip with samples of different geographical origin.

Table 7: HIV-1 protease wild-type and drug-selected mutation probes with their corresponding sequences as applied on the HIV-1 protease LiPA strip. The most frequently observed wild-type sequence is shown at the top line. Probe names corresponding to the selected motifs are indicated in the left column, the relevant part of each probe applied on the strip is shown under the consensus sequence.

#### **EXAMPLES**

#### 30 Example 1:

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## Selection of the plasma samples, PCR amplification and cloning of the PCR products.

<u>Plasma samples</u> (n=557) were taken from HIV type-1 infected patients and stored at -20°C until use. Plasma samples were obtained from naive and drug-treated patients. The drugs involved ritonavir, indinavir and saquinavir. The serum samples were collected from patients residing in Europe (Belgium, Luxembourg, France, Spain and UK), USA and Brazil.

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HIV RNA was prepared from these samples using the guanidinium-phenol procedure. Fifty μl plasma was mixed with 150 μl Trizol®LS Reagent (Life Technologies, Gent, Belgium) at room temperature (volume ratio: 1unit sample/ 3 units Trizol ). Lysis and denaturation occurred by carefully pipetting up and down several times, followed by an incubation step at room temperature for at least 5 minutes. Fourthy μl CHCl<sub>3</sub> was added and the mixture was shaken vigorously by hand for at least 15 seconds, and incubated for 15 minutes at room temperature. The samples were centrifuged at maximum 12,000g for 15 minutes at 4°C, and the colorless aqueous phase was collected and mixed with 100 μl isopropanol. To visualize the minute amounts of viral RNA, 20 μl of 1μg/μl Dextran T500 (Pharmacia) was added, mixed and left at room temperature for 10 minutes. Following centrifugation at max. 12,000g for 10 minutes at 4°C and aspiration of the supernatant, the RNA pellet was washed with 200 μl ethanol, mixed by vortexing and collected by centrifugation at 7,500g for 5 minutes at 4°C. Finally the RNA pellet was briefly air-dried and stored at -20°C. Alternatively, the High Pure Viral Nucleic Acid Kit (Boehringer Mannheim) was used to extract RNA from the samples

For cDNA synthesis and PCR amplification, the RNA pellet was dissolved in 15  $\mu$ l random primers (20 ng/ $\mu$ l, pdN<sub>6</sub>, Pharmacia), prepared in DEPC-treated or HPLC grade water. After denaturation at 70°C for 10 minutes, 5  $\mu$ l cDNA mix was added, composed of 4  $\mu$ l 5x AMV-RT buffer (250mM Tris.HCl pH 8.5, 100mM KCl, 30mM MgCl<sub>2</sub>, 25 mM DTT), 0.4  $\mu$ L 25mM dXTPs, 0.2  $\mu$ l or 25U Ribonuclease Inhibitor (HPRI, Amersham), and 0.3  $\mu$ l or 8U AMV-RT (Stratagene). cDNA synthesis occurred during the 90 minutes incubation at 42°C. The HIV -1 protease gene was than amplified using the following reaction mixture: 5  $\mu$ l cDNA, 4.5  $\mu$ l 10x Taq buffer, 0.3  $\mu$ l 25 mM dXTPs, 1  $\mu$ l (10 pmol) of each PCR primer, 38  $\mu$ l H<sub>2</sub>O, and 0.2  $\mu$ l (1 U) Taq. . Alternatively, the Titon One Tube RT-PCR system (Boehringer Mannheim) was used to perform RT-PCR.

Codon positions involving resistance to saquinavir, ritonavir, indinavir, nelfinavir and VX-478 have been described (Shinazi et al) and PCR amplification primers were chosen outside these regions. The primer design was based on HIV-1 published sequences (mainly genotype B clade) (Myers et al.) and located in regions that showed a high degree of nucleotide conservation between the different HIV-1 clades. The final amplified region covered the HIV-1 protease gene from codon 9 to codon 99. The primers for biosequence: outer sense primer Pr16: following the amplification had CAGAGCCAACAGCCCACCAG3' (SEQ ID NO 1); nested sense primer Prot 2 bio: 5' CCT CAR ATC ACT CTT TGG CAA CG 3' (SEQ ID NO 3); nested antisense primer Prot 6 bio: 3' TAA TCR GGA TAA CTY TGA CAT GGT C 5' (SEQ ID NO 4); and outer antisense primer RT12: 5' bioATCAGGATGGAGTTCATAACCCATCCA3' (SEQ ID NO 2). Annealing occurred at 57°C, extension at 72°C and denaturation at 94°C. Each step of the cycle took 1 minute, the outer PCR contained 40 cycles, the nested round 35. Nested round PCR products were analyzed on agarose gel and only clearly visible amplification products were used in the LiPA procedure. Quantification of viral

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RNA was obtained with the HIV Monitor<sup>TM</sup>test (Roche, Brussels, Belgium). Later on, new sets of primers for amplification were selected. For the amplification of HIV protease codon 30-84: outer sense primer prot16: 5'-CAGAGCCAACAGCCCCACCAG-3' (SEQ ID NO 501), outer antisense primer prot5: 5'-TTTTCTTCTGTCAATGGCCATTGTTT-3' (SEQ ID NO 502) were used. Annealing occurred at 50°C, extension at 68°C and denaturation at 94°C for 35 cycles for the outer PCR. For the nested PCR annealing occurred at 45°C, denaturation at 94°C and extension at 92°C with primers: nested sense primers prot2a-bio: 5'-bio-CCTCAAATCACTCTTTGGCAACG-3' (SEQ ID NO 503) and prot2b-bio: 5'-bio-CCTCAGATCACTCTTTGGCAACG-3' (SEQ ID NO 504), and nested antisense primer prot31bio: 5'-bio-AGTCAACAGATTTCTTCCAAT-3' (SEQ ID NO 6). For the amplification of HIV protease codon 90, the outer PCR was as specified for HIV protease codon 30-84. For the nested PCR, nested sense primer prot41-bio: 5'-bio-CCTGTCAACATAATTGCAAG-3' (SEQ ID NO 505) and nested antisense primers prot6a: 5'-bio-CTGGTACAGTTTCAATAGGGCTAAT-3' (SEQ ID NO 506), prot6b: 5'-bio-5'-bio-CTGGTACAGTTTCAATAGGACTAAT-3' (SEO ID NO 507), prot6c: CTGGTACAGTCTCAATAGGACTAAT-3' (SEO IDNO 508), prot6d: 5'-bio-CTGGTACAGTCTCAATAGGGCTAAT-3' (SEQ ID NO 509) were used. For the nested PCR the annealing temperature occurred at 45°C. Primers were tested on a plasmid, which contained an HIV fragment of 1301 bp ligated in a pGEM-T vector. The fragment contains protease, reverse transcriptase and the primer sites of first and second round PCR. By restriction with Sac I the plasmid is linearised.

Selected PCR products were cloned into the pretreated EcoRV site of the pGEMT vector (Promega). Recombinant clones were selected after α-complementation and restriction fragment length analysis, and sequenced using standard sequencing techniques with plasmid primers and internal HIV protease primers. Sometimes biotinylated fragments were directly sequenced with a dye-terminator protocol (Applied Biosystems) using the amplification primers. Alternatively, nested PCR was carried out with analogs of the nested primers, in which the biotin group was replaced with the T7- and SP6-primer sequence, respectively. These amplicons were than sequenced with an SP6- and T7-dye-primer procedure.

#### Example 2:

#### 30 Selection of a reference panel

Codon positions involving resistance to saquinavir, ritonavir, indinavir, nelfinavir and VX-478 have been described (Shinazi *et al. 1997*). It was the aim to clone in plasmids those viral protease genes that are covering the different genetic motifs at those important codon positions conferring resistance against the described protease inhibitors.

After careful analysis of 312 protease gene sequences, obtained after direct sequencing of PCR fragments, a selection of 47 PCR fragments which covered the different target polymorphisms and

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mutations were retained and cloned in plasmids using described cloning techniques. The selection of samples originated from naive or drug-treated European, Brazilian or US patients. These 47 recombinant plasmids are used as a reference panel, a panel that was sequenced on both strands, and biotinylated PCR products from this panel were used to optimize probes for specificity and sensitivity.

Although this panel of 47 samples is a representative selection of clones at this moment, it is important to mention here that this selection is an fact only a temporally picture of the variability of the virus, and a continuous update of this panel will be mandatory. This includes on ongoing screening for the new variants of the virus, and recombinant cloning of these new motifs.

#### Probe selection and LiPA testing.

To cover all the different genetic motifs in the reference panel, a total of 471 probes were designed (codon 30: 40 probes; codon 46/48: 72 probes; codon 50:55 probes; codon 54: 54 probes; codon 82/84: 130 probes; codon 90: 120 probes). Table 3 shows the different probes that were selected for the different codon positions.

It was the aim to adapt all probes to react specifically under the same hybridization and wash conditions by carefully considering the % (G+C), the probe length, the final concentration of the buffer components, and hybridization temperature (Stuyver et al., 1997). Therefore, probes were provided enzymatically with a poly-T-tail using the TdT (Pharmacia) in a standard reaction condition, and purified via precipitation. For a limited number of probes with 3' T-ending sequences, an additional G was incorporated between the probe sequence and the poly-T-tail in order to limit the hybridizing part to the specific probe sequence and to exclude hybridization with the tail sequence. Probe pellets were dissolved in standard saline citrate (SSC) buffer and applied as horizontal parallel lines on a membrane strip. Control lines for amplification (probe 5' TAGGGGGAATTGGAGGTTTTAG 3', HIV protease aa 47 to aa 54) and conjugate incubation (biotinylated DNA) were applied alongside. Probes were immobilized onto membranes by baking, and the membranes were sliced into 4mm strips also called LiPA strips.

Selection of the amplification primers and PCR amplification was as described in example 1. In order to select specific reacting probes out of the 471 candidate probes, LiPA tests were performed with biotinylated PCR fragments from the reference panel. To perform LiPA tests, equal amounts (10 µl) of biotinylated amplification products and denaturation mixture (0.4 N NaOH/0.1% SDS) were mixed, followed by an incubation at room temperature for 5 minutes. Following this denaturation step, 2 ml hybridization buffer (2xSSC, 0.1% SDS, 50mM Tris pH7.5) was added together with a membrane strip and hybridization was carried out at 39°C for 30 min. Then, the hybridization mixture was replaced by stringent washing buffer (same composition as hybridization buffer), and stringent washing occurred first at room temperature for 5 minutes and than at 39°C for another 25 minutes. Buffers were than replaced to be suitable for the streptavidine alkaline phosphatase conjugate incubations. After 30 minutes

incubation at room temperature, conjugate was rinsed away and replaced by the substrate components for alkaline phosphatase, Nitro-Blue-Tetrazolium and 5-Bromo-4-Chloro-3-Indolyl Phosphate. After 30 minutes incubation at room temperature, probes where hybridization occurred became visible because of the purple brown precipitate at these positions.

After careful analysis of the 471 probes, the most specific and sensitive probes (n=46) were finally selected, covering the natural and drug-selected variability in the vicinity of aa. 30, 46, 48, 50, 54, 82, 84, and 90. Figure 2 shows the reactivity of the finally selected probes with the reference panel.

### Example 3:

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#### LiPA testing on clinical samples.

A total of 856 samples were tested on this selection of 46 specific probes. The geographical origin of these samples is as follows: USA:359; France: 154; UK:36; Brazil 58; Spain 35; Belgium 199; Luxembourg: 15.

From this population, a total of 144 samples were sequenced which allowed to separate the genotype B samples (94) from the non-B samples (50). After analysis of these genotyped samples on LiPA, the genotypic reactivity on the selected probes was scored. Figures 4A to 4F show these results for the different codon positions and for the genotype B versus non-B group. From these tables, it is clear that there is little difference in sequence usage for the different codon positions with respect to specific reactivities at the different probes.

The total collection of 856 samples was then tested on the available 46 probes. After dissection of these reactivities over the different probes and different geographical origin, the picture looks as is presented in Figures 5A to 5F. Again here, the majority of the sequences used at the different codon positions are restricted to some very abundant wild type motifs. It is important to mention here that the majority of these samples are taken from patients never treated with protease inhibitors, en therefore, the majority of the reactivities are found in wild type motifs. Nevertheless, it is clear from some codon positions that the variability at some codon positions in the mutant motif might be considerable, and again, a continuos update on heavily treated patients is mandatory. Another issue is the amount of double blank reactivities, which is in this approach reaching up to 5% in global; with some peak values for some countries for some codon positions: for example 13.8% for codon 82/85 in Brazil; and 18.1 % for codon 90 in Belgium.

The continuous update resulted in a further selection of probes. This later configuration of the strip is indicated in table 7.

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# Table 1

pc30w25 pc30w29 pc30w32 pc30w36	26 27 28 29 30 31 32 33 34 ACA GGA GCA GAT GAT ACA GTA TTA GAA G GCA GAT GAT ACA GT A GCG GAT GAT ACA GCA GAT GAC ACA GT GCA GAC GAT ACA GG A GCA GAT AAT ACA GT		14 13 14 14 14	Seq ID 31 35 38 42 29
pc48w47 pc48w45 pc48w72 pc48w41	44 45 46 47 48 49 50 51 52 CCA AAA ATG ATA GGG GGA ATT GGA GGT AAA ATG ATA GGG GGA A ATG ATA GGA GGA ATT A AAA ATA ATA GGG GGA ATG ATA GGG GGA ATG ATA GGG GGA ATG ATA GTG GGA ATT	42 42 42 40	15 16 16 15	93 91 120 87
pc50w31 pc50w44 pc50w52 pc50m37	48 49 50 51 52 53 54 GGG GGA ATT GGA GGT TTT ATC GGA ATT GGA GGT TTT GGA ATT GGG GGT TTG GA ATT GGA GGC TTG GGG GGA GTT GGA	42 42 40	15 15 14 12	151 164 172 157
pc54w3 pc54w34 pc54w14 pc54w19 pc54w22 pc54w26 pc54w27 pc54w55 pc54m35 pc54m37	51 52 53 54 55 56 57 58  GGA GGT TTT ATC AAA GTA AGA CAG GT TTT ATC AAA GTA AGA GA GGT TTT ATC AAA GT GGT TTT ATC AAA GT A GGC TTT ATC AAA GT A GGC TTT ATC AAA GT A GGT TTT ATT AAA GTA A GGT TTC ATT AAA GTA A GGT TTC ATT AAA GTA A GGT TTT ATT AAA GTA GGT TTT GCC AAA GT GGT TTT GCC AAA GT GGT TTT GTC AAA GTA GGT TTT GTC AAA GTA	42 42 42 42 42 40 38 40 42	17 16 16 16 17 16 16 15 15	178 212 189 194 197 202 204 213 215
pc82w91 pc82w60 pc82w111 pc82w89 pc82w42 pc82m36 pc82m67 pc82m38 pc82m10 pc82m10 pc82m12	ACA CCT GTT AAC ATA AG CA CCT GTC AAC GTA ACA CCT ACC AAC ATA ACA CCT ACC AAC GT ACA CCT TTC AAC ATA ACA CCT TTC AAC ATA CA CCT TTC AAC ATA CA CCT GCC AAC ATA CA CCT GCC AAT ATA	87 AGA 44 42 44 42 42 40 44 44 44 44	17 16 17 14 15 14 15 15 17 15 16	318 287 338 316 269 263 294 265 332 354 267 290 328

# Table 1 - Cont'd

	86	87	88	89	90	91	92	93	94			
	GGA	AGA	AAT	CTG	TTG	ACT	CAG	ATT	GGT			
pc90w27			AAT	CTG	TTG	ACT	CA			38	14	384
pc90w37			AAT	CTG	TTG	ACT	CAG	ATG		42	18	394
pc90w39		GA	ACT	CTG	TTG	ACT	C			44	15	396
pc90w50			AAT	ATG	TTG	ACT	CAG			40	15	407
pc90w52			AAT	TTG	TTG	ACT	CAG			40	15	409
pc90w69		GA	AAC	CTG	TTG	ACT				40	14	426
pc90w73				ΤG	TTG	ACA	CAG	CTT	G	44	15	430
pc90w79				TG	TTG	ACC	CAG	ATT	G	44	15	436
pc90m43		A	AAT	CTG	ATG	ACT	CA			40	15	400
pc90m56			AAT	ATG	ATG	ACC	CAG			42	15	413

Table 2
Protease Inhibitors

Compound	Amino acid change	Codon change
Protease Inhibitors		
A-77003	R8Q R8K V32I	CGA to CAA CGA to AAA GTA to ATA
	M46I	ATG to ATA
	M46L M46F M46V G48V A71V	ATG to TTC ATG to TTC ATG to GTG GGG to GTG GCT to GTT
	V82I	GTC to ATC
	V82A	GTC to GCC
	L63P A71T A71V G73S V82A V82F V82T I84V	CTC to CCC GCT to ACT GCT to GTT GGT to GCC GTC to TTC GTC to ACC ATA to GTA
	L90M	TTG to ATG
P9941	V82A	GTC to GCC
Ro 31-8959 (saquinavir)	L10I G48V	CTC to ATC GGG to GTG
	I54V I54V G73S V82A I84V L90M	ATC to GTC ATA to GTA GGT to AGT GTC to GCC ATA to GTA TTG to ATG
RPI-312	I84V	ATA to GTA

29 **Table 2 - Cont'd-1** 

SC-52151	L24V G48V A71V	TTA to GTA GGG to GTG GCT to GTT
	V75I P81T V82A N88D	GTA to ATA CCT to ACT GTC to GCC AAT to GAT
SC-55389A	L10F N88S	CTC to CGC AAT to AGT
SKF108842	V82T I84V	GTC to ACC ATA to GTA
SKF108922	V82A V82T	GTC to GCC GTC to ACC
VB 11,328	L10F M46I I47V I50V 184V	CTC to GGC ATG to ATA ATA to CTA ATT to GTT ATA to GTA
VX-478 (141W94)	L10F M46I I47V I50V I84V	CTC to CGC ATG to ATA ATA to CTA ATT to GTT ATA to GTA
XM323	L10F K45I M46L V82A	CTC to CGC AAA to ATA ATG to CTG GTC to GCC
	V82I V82F I84V	GTC to ATC GTC to TTC ATA to GTA
	L97V	TTA to GTA
	I82T	ATC to ACC
A-75925 ABT-538 (ritonavir)	V32I K20R L33F	GTA to ATA AAG to AAA TTA to TTC

Table 2 - Cont'd-2

	M36I	ATG to ATA
	M46I	ATG to ATA
	I54L	ATC to?
	I54V	ATC to GTC
	A71V	GTC to GTT
	V82F	GTC to TTC
	V82A	GTC to GCC
	V82T	GTC to ACC
	V82S	GTC to TCC
	I84V	ATA to GTA
	L90M	TTG to ATG
AG1343		
(nelfinavir)	D30N	GAT to AAT
(	M36I	
	M46I	ATG to ATA
	L63P	CTC to CCC
	A71V	GCT to GTT
	V771	
	184V	ATA to GTA
	N88D	
	L90M	TTG to ATG
BILA 1906	V32I	GTA to ATA
BS	M46I	ATG to ATA
	M46L	ATG to TTG
	A71V	GCT to GTT
	I84A	ATA to GCA
	184V	ATA to GTA
BILA 2011	V32I	GTA to ATA
(palinavir)	A71V	GCT to GTT
	I84A	ATG to ATA
	L63P	CTC to CCC
BILA 2185 BS	L23I	CTA to ATA
BMS 186,318	A71T	GCT to ACT
210,310	V82A	GTC to GCC
DMP 450	L10F	CTC to TTC

Table 2 - Cont'd-3

	M46I D60E I84V	ATG to ATA GAT to GAA ATA to GTA
KNI-272	V32I	GTA to ATA
MK-639 (L-735,524, indinavir)	L10I L10R L10V K20M K20R L24I V32I M46I M46L I54V	CTC to ATC CTC to CGC CTC to GTC AAG to ATG AAG to AAA TTA to ATA GTA to ATA ATG to ATA ATG to TTG ATC to GTC

# Table 3

	26	27	28	29	30	31	32	33	34	35	length	Seq ID
	ACA	GGA				ACA			GAA	GAA	18	7
P30w1			GCA			ACA		TT			18 19	8
P30w2			GCA			ACA		TT			19	9
P30w3			GCA			ACA		TTA			20	10
P30w4			GCA			ACA					21	11
P30w5	7 (7		GCA			ACA	GIA	IIA			18	12
P30w6		GGA				ACA	CIT				19	13
P30w7		GGA				ACA		m.c			20	14
P30w8	А	GGA				ACA ACA					19	15
P30w9	70 70		GCA			ACA		10			19	16
P30w10	ACA	GGA				ACA		an an			18	17
P30m11			GCA GCA		AAT		GTA				19	18
P30m12			GCA								19	19
P30m13			GCA			ACA					20	20
P30m14			GCA			ACA					21	21
P30m15	7\ (~7\	GGA			AAT	ACA	GIA	TIL			18	22
P30m15		GGA				ACA	СT				19	23
P30m17		GGA				ACA		TС			20	24
P30m18 P30m19	A		GCA		AAT		GTA				19	25
P30m19	$\mathcal{N} \subset \mathcal{N}$	GGA			AAT	ACA		10			19	26
p30m20	ACA		GCA		GAT	ACA					15	27
p30w21			GCA		GAT		GTA	G			16	28
p30w22 p30m23			GCA		AAT		GTA	J			15	29
p30m24			GCA		AAT		GTA	G			16	30
p30w25					GAT						14	31
p30w26		A	GCA		GAT						14	32
p30w27				GAT	GAT	ACA					13	33
p30w28		GA	GCG			ACA					14	34
p30w29			GCG		GAT	ACA					13	35
p30m30				GAT			GTA				15	36
p30m31			GCA	GAT	AAT	ACA	GT				14	37
p30w32				GAT			GT				14	38
p30w33			CA	GAT	GAC	ACA	GTA	G			14	39
p30w34			CA	GAT	GAT	ACA	ATA	${ m TT}$			16	40
p30w35			GCA	GAT	GAT	ACA	ATA	TG			16	41
p30w36			GCA	GAC	GAT	ACA	GG				13	42
p30w37			GCA	GAC							14	43
p30w38			A	GAT		ACA					15	44
p30w39			Α				ATA		7		16	45
p30w40			GCA	GAT	GAT	ACA	ATA				15	46

Table 3 - Cont'd-1

	44	45	46	47	48	49	50	51	52	53	54	length	Seq ID
		AAA			GGG	GGA	ATT		GGT	TTT	ATC	_	
P48w1	00					GGA		GGA		GG		18	47
P48w2						GGA		GGA		TG		19	48
P48w3						GGA		GGA		TTG		20	49
						GGA		GGA		TTT		21	50
P48w4			G			GGA			GGT			21	51
P48w5			_			GGA		GGA	001	110		18	52
P48w6			ATG						<u></u>			19	53
P48w7		_				GGA		GGA	G			19	54
P48w8						GGA		GGA	~				55
P48w9						GGA		GGA		~~		20	
P48w10		Α				GGA			GGG	GG		22	56
P48w21						GGA		GGA				18	57
P48w22						GGA		GGA				18	58
P48w23						GGA		GGA				19	59
P48w24		А	ATG			GGA		GGA				19	60
P48w25						GGA			GGT			18	61
P48w26						GGA		GGA		ΤG		19	62
P48w28						GGA			GGT	TTG		20	63
P48w29				ATA	GGG	GGA	ATT	GGA	GGT	TTT		21	64
P48m11				GTA	GTG	GGA	ATT	GGA	GGT	GG		18	65
P48m12				GTA	GTG	GGA	ATT	GGA	GGT	TG		19	66
P48m13				GTA	GTG	GGA	ATT	GGA	GGT	TTG		20	67
P48m14				GTA	GTG	GGA	ATT	GGA	GGT	TTT		21	68
P48m15			G	GTA	GTG	GGA	ATT	GGA	GGT	TTG		21	69
P48m16			ATG			GGA		GGA				18	70
P48m17						GGA		GGA	G			19	71
P48m18		<b>Z</b> A				GGA		GGA				19	72
P48m19							ATT					20	73
P48m20							ATT		GGG	GG		22	74
P48m29		7.	2120				ATT		GGT			18	75
P48m30						GGA			GGT			19	76
P48m31			ATG				ATT					18	77
P48m32							ATT					19	78
P48m33		7\					ATT					19	79
p48w34		73					ATT		-			14	80
p48w34 p48w35							ATT					15	81
							ATT					16	82
p48w36						GGA		00				15	83
p48w37							ATT	G				14	84
p48m38							ATT					15	85
p48m39												16	86
p48m40							ATT					15	87
p48m41							TTA						88
p48w42							ATT					15	
p48w43							GTT					14	89
p48w44		_					GTT					14	90 01
p48w45		P					ATT					16	91
p48w46							LTA 1	•				15	92
p48w47						GGA	7					15	93
p48w48	I	AAA A	ATC	ATA	GGG	GG						15	94

## Table 3 - Cont'd-2

p48w49		AA	ATG	ATA	GGG	GGA	AG			15	
p48w50		AAA	ATA	ATA	GGG	GGA	AG			16	
p48w51		AAA	ATA	AAA	AT					15	
p48m52		AAA	ATG	ATA	GTG	GGA	AG			16	
p48w52b		AAA		ATA		GG				14	
p48m53		AAA	ATG	ATA		GGA				15	
p48w53b		AAA		ATA		GGA				15	
p48w54	CA	AAA		ATA	G					15	
p48w55			ATG	GTA		GGA	ATT			15	
p48w56		AΑ	ATG	GTA	GGG	GGA				14	
p48w57	A	AAA	ATG	GTA	GGG	G	7 mm			14 15	
p48w58			ATG	ATA	GGG	GAA		~~ 7		15	
p48w59				ATA	GGG	GAA		GGA	C	16	
p48w60				ATA	GGG	GAA	ATT	GGA	G	15	
p48w61			ATG	ATA	GGG	GGG	ATT	CC		14	
p48w62				ATA	GGG	GGG	ATT	GG		13	
p48w63			7. 55. 7	A	GGG	GGG	TTA	GGA		1:	
p48m64	-		ATA			GGA				10	
p48m65	A			ATA	GTG GTG	GGA GG				1	
p48m66	CA		ATA			GGA				1	
p48m67	71	AAA	TTG TTG	ATA ATA						1	16
p48m68	A	AAA								1	17
p48m69	CA	AAA AAA								1	18
p48w70	7\		ATG							1	 19
p48w71 pc48w72	A A		ATA							1	 20
DCGOWIZ	17	1 77 75 7	*****								

Table 3 - Cont'd-3

	45	46	47	48	49	50	51	52	53	54	length	Seq ID
	AAA	ATG	GTA	GGG	GGA	ATT	GGA	GGT	TTT	ATC		
P50w1				GGG	GGA	TTA	GGA	GGT	TTT		18	121
P50w2			A	GGG	GGA	ATT	GGA	GGT	TTT		19	122
P50w3			TA	GGG	GGA	ATT	GGA	GGT	TTT		20	123
P50w4			A	GGG	GGA	ATT	GGA	GGT	TTT	AG	20	124
P50w5			TA	GGG	GGA	ATT	GGA	GGT	TTT	AG	21	125
P50w6			GTA	GGG	GGA	ATT	GGA	GGT	TGG		19	126
P50w7		G	GTA	GGG	GGA	ATT	GGA	GGT	TGG		20	127
P50w8			GTA	GGG	GGA	ATT	GGA	GGT	TTG		20	128
P50w9			GTA	GGG	GGA	ATT	GGA	GGT	TTT		20	129
P50w10		TG	GTA	GGG	GGA	ATT	GGA	GGT	GG		20	130
p50w21				GG	GGA	ATT	GGA	GGT	TTT		17	131
P50w22				GG	GGA	ATT	GGA	GGT	TTG		16	132
P50w23				GG	GGA	ATT	GGA	GGT	TTT	AG	18	133
P50w24				GG	GGA	ATT	GGA	GGT	TG		15	134
P50w25				G	GGA	ATT	GGA	GGT	TTT	AT	18	135
P50w26				GG	GGA	ATT	GGA	GGT	TTT		17	136
P50m11				GGG	GGA	GTT	GGA	GGT	TTT		18	137
P50m12			А	GGG	GGA	GTT	GGA	GGT	TTT		19	138
P50m13			TA	GGG	GGA	GTT	GGA	GGT	TTT		20	139
P50m14			А	GGG	GGA	GTT	GGA	GGT	TTT	AG	20	140
P50m15			ΤA	GGG	GGA	GTT	GGA	GGT	TTT	AG	21	141
P50m16			GTA	GGG	GGA	GTT	GGA	GGT	' TGG		19	142
P50m17		(	GTA	GGG	GGA	GTT	GGA	GGT	' TGG		20	143
P50m18			GTA	GGG	GGA	GTT	GGA	GGI	TTG		20	144
P50m19			GTA	GGG	GGA	GTT	GGA	GGT	TTT	ATC	21	145
P50m20		TO	GTA	GGG	GGA	GTT	GGA	GGI	' GG		20	146
P50m27				GG	GGA	GTT	GGA	GGI	TTG	;	19	147
P50m28				GG	GGA	GTI	GGA	GGI	TTT	'AG	18	148
P50m29				GG	GGA	GTT	GGA	GG1			15	149
P50m30				G	GGA	GTI	GGA	GG'	TTT	TA :	18	150
p50w31					GGF	TTA A	GGF	GGT	rtt 1	-	15	151
p50w32				0	GGF	TTA A	GGF	A GGT	r TGC	3	15	152
p50m33					GGF	A GTT	GGF	A GG	r TTI	2	15	153
p50m34					G GGF	GTI	GGA	A GG	r TGC	,	14	154
p50m35				GGC	G GGI	A GT	GG <i>I</i>	4 G			13	155
p50m36						A GTT					12	156
p50m37				GG			r GGZ				12	157
p50w38							r gg				14	158
p50w39					GZ	A AT	r GG	G GG'	r TT	Γ	14	159

## Table 3 - Cont'd-4

p50w40		GΑ	ATT	GGG	GGT	TTT AG	15	160
p50w40		GGA	ATT	GGG	GGT	TG	13	161
p50w41					GGT		12	162
p50w42		GA	ATT	GGG	GGT	TG	12	163
p50w45		GA	ATT	GGG	GGT	TTG	13	
p50w45	GGG	GGA	ATT	GCA	G		13	165
p50w45		GGA	ATT	GCA	GGT	TG	14	166
p50w47		GGA	ATT	GCA	GGT	G	13	167
p50w47		GGA	ATT	GGA	GGG	TTG	14	168
p50w49		GΑ	ATT	GGA	GGG	TTG	13	3 169
p50w50		GΑ	ATT	GGA	GGG	TTT	14	170
p50w51		GGA	ATT	GGA	GGC	TTG	14	1 171
p50w51		GΑ	ATT	GGA	GGC	TTG	13	3 172
=		GΑ	ATT	GGA	GGC	TTT	14	173
p50w53		GGA			GGT	TTG	15	5 174
p50m54 p50m55		GA			GGT		1	4 175
P00:::00								

Table 3 - Cont'd-5

		52	53	54	55	56	57	58	length	Seq ID
F 4 1				ATC	AAA	GTA GTA		CAG	16	176
p54w1		GGI	TTT		AAA				16	177
p54w2		GT	$\mathrm{TTT}$		AAA				17	178
p54w3 p54w4		T	TTT		AAA				16	179
p54w4 p54w5		GGT	TTT		AAA				15	180
p54w6		GT	TTT		AAA				15	181
p54m7		GGT	TTT		AAA				15	182
p54m8		GT	TTT		AAA		A		15	183
p54m9		GT	TTT		AAA				16	184
p54m10		T	TTT		AAA				16	185
p54m11		GGT	TTT	GCC	AAA	GT			14	186
p54m12		GT	TTT	GCC	AAA	GTA			14	187
p54w13		GT	TTT	ATC	AAG	GTA	AA		16	188
p54w14		GGT	TTT		AAG		A		16	189
p54w15	A	GGT	TTT		AAG				16	190
p54w16		GT	TTT	ATC		GTC			17	191
p54w17			TTT		AAA			С	16	192
p54w18	A	GGC	TTT		AAA		A		17	193
p54w19	A	GGC		ATC		GTA			16	194
p54m20	A	GGT	TTT	ATT		GTA			17	195
p54m21		GGT	TTT	ATT		GTA	AG		17	196
p54w22	GA		TTT	ATT					17	197
p54m22	GA		TTT	ATT					17	198
p54m23		GGT	TTT	ATT					16 15	199 200
p54m24		GGT	TTC	ATT		GTA			16	201
p54m25	_	GGT	TTC	ATT					16	202
p54w26		GGT	TTC	ATT					16	203
p54m26	А	GGT	TTC						16	204
p54w27		GGT	TTT	TTA TTA					16	205
p54m27	7\	GGT	TTT TTT						16	206
p54m28		GGT	TTT						16	207
p54m29	GA						AG		17	208
p54m30		GGT GGT			CAAA				16	209
p54w31	71	GGT			AAA				17	210
p54w32 p54w33		GGT			C AAA				16	211
p54w34		GGT			CAAA		_		16	212
p54m35	Q23	GGT			CAAA		Ā		15	213
p54m36		GGT			CAAA				16	214
p54m37		GGI			CAGA				15	215
p54m38		GGT				A GTA			16	216
p54w39		GGG			CAA	A GTA	Ā		15	217
p54w40		GGG		TA	CAAA		$A \in A$		16	218
p54w41		GGC		CAT	CAAA	A GT			14	219
p54w42	G <i>P</i>	GGC	TTC		CAA				14	220
p54m48		GGT			C AA				14	221
p54m49		GI	TT	r GT	C AG	A GT	Ą		14	222

### Table 3 - Cont'd-6

. F 4 F O		CCIT	արդուր	CTC	AGA	CT	14	223
p54m50								
p54w51	A	GGT	TTA	ATC	AAA	GTA	16	224
p54w52	GA	GGT	TTA	ATC	AAA	GT	16	225
p54m53		GGT	TTT	ACC	AAA	GTA	15	226
p54m54		GGT	TTT	ACC	AAA	GT	14	227

Table 3 - Cont'd-7

	78	79	80	81	82	83	84	85	86	87	length	Seq	ID
P82w1 P82w2 P82w3 P82w4 P82w5	A A	CCT CCT CCT CCT	ACA ACA ACA ACA ACA	CCT CCT CCT CCT	GTC GTC GTC GTC	AAC AAC AAC AAC	ATA ATA ATA ATA	AG ATG ATT AG	GGA	AGA	19 20 21 20 21	22 22 23 23 23	9 0 1 2
P82w6 P82w7 P82w8 P82w9 P82w10 P82W21			A ACA	CCT CCT CCT	GTC GTC GTC GTC	AAC AAC AAC		ATT ATT	GGA GG	A	19 20 20 20 20 19	23 23 23 23 23 23	4 5 6 7 8
P82m11 P82m12 P82m13 P82m14 P82m15	A	CCT CCT CCT	ACA ACA ACA ACA	CCT CCT CCT	ACC ACC ACC ACC	AAC AAC AAC AAC	ATA ATA ATA ATA	ATG ATT AG			19 20 21 20 21	23 24 24 24 24	0 1 2 3
P82m16 P82m17 P82m18 P82m19 P82m20 P82m22		CCT	A	CCT CCT CCT	ACC ACC ACC ACC	AAC AAC AAC	ATA ATA ATA	ATT ATT			19 20 20 20 19 21	24 24 24 24 24 24	5 6 7 8
P82m23 P82m24 P82m25 P82m26 P82m27		CCT	ACA ACA A A	CCT CCT	GCC TCC TTC GCC	AAC AAC AAC AAC	ATA	ATT ATT ATT ATT	GGA GGA GGA	A	21 21 20 20 20	25 25 25 25 25	0 1 2 3
P82m28 P82m29 P82m30 P82m31 P82w32			A A A ACA	CCT CCT CCT	TTC GCC TCC GTC	AAC AAC AAC		ATT ATT	GGA		16 19 19 15	25 25 25 25 25	66 57 58 59
P82w33 P82w34 P82w35 P82m36 P82m37		Т	ACA CA ACA CA	CCT CCT CCT	GTC GTC ACC ACC	AAC AAC AAC AAC	ATA ATA ATA				16 15 14 15	26 26 26 26	51 52 53 54
P82m38 P82m39 P82m40 P82m41 P82w42			CA ACA CA CA	CCT CCT CCT	TTC GCC GCC GTC	AAC AAC AAC AAC AAC	ATA ATA ATA GTA				15 14 15 14 14	26 26 26 26 26	56 57 58 59
P82w43 P82w44 P82w45 P82w46 P82m47		Т	ACA ACG ACG	CCT CCT CCT	GTC GTC GTC	AAC AAC AAC	AT				15 15 15 15	27 27 27	

# Table 3 - Cont'd-8

D0040		14 275
P82m48 P82m49	CA CCT TCC AAC ATA ACA CCT TCC AAC AT	14 275 14 276
P82m50	ACA CCT ATC AAC ATA	15 277
P82m51	CA CCT ATC AAC ATA AG	15 278
P82m52	CA CCT ATC AAC ATA ATG	16 279
P82m53	A CCT ATC AAC ATA ATG	15 280
P82w54	CCT GTC AAC ATA ATT	15 281
P82w55	CCT GTT AAC ATA ATT G	16 282
P82w56	A CCT GTT AAC ATA ATG	15 283
P82w57	CCG GTC AAC ATA ATT	15 284
P82w58	ACG CCT GTC AAC AT	14 285
P82w59	CCT GTC AAT ATA ATT	15 286
P82w60	CA CCT GTC AAT ATA ATG	16 287
P82w61	ACA CCT GTC AAT ATA AG	16 288
P82m62	CCT GCC AAT ATA ATT	15 289
P82m63	CA CCT GCC AAT ATA AG	15 290
P82m64	CCT ACC AAC GTA ATT	15 291
P82m65	CCT ACC AAC GTA ATG	14 292
P82m66	CA CCT ACC AAC GTA	14 293
P82m67	ACA CCT ACC AAC GT	14 294
P82m68	CCT TTC AAC GTA ATT	15 295
P82m69	CA CCT TTC AAC GTA AG	15 296 15 297
P82m70	ACA CCT TTC AAC GTA	15 297 15 298
P82m71	A CCT TTC AAC GTA ATG CT GTC AAT ATA ATT G	15 296
p82w72	CT GTC AAT ATA ATT G CCT GTC AAT ATA ATT G	16 300
p82w73 p82w74	A CCT GTC AAT ATA ATT	16 301
p82w74 p82w75	CT GTC AAT ATA ATT GG	16 302
p82w75 p82w76	CCT ACG CCT GTC AA	14 303
p82w77	CT ACG CCT GTC AAC	14 304
p82w78	A CCT ACG CCT GTC AA	15 305
p82w79	A CCT ACG CCT GTC A	14 306
p82w80	T ACA CCG GTC AAC A	14 307
p82w81	CT ACA CCG GTC AA	13 308
p82w82	CCT ACA CCG GTC A	13 309
p82w83	CA CCT GTC AAC ATA A	15 310
p82w84	A CCT GTC AAC ATA AT	15 311
p82w85	CT ACA CCT GTC AAC A	15 312
p82w86	ACA CCT GTC AAC AT	14 313
p82w87	A CCT GTT AAC ATA ATT G	17 314
p82w88	CA CCT GTT AAC ATA AG	15 315
p82w89	ACA CCT GTT AAC ATA AG	16 316
p82w90	TCA CCT GTC AAC ATA	14 317
p82w91	ACA CCT GTC AAC ATA A	16 318 16 319
p82w92	CA CCT GTC AAC ATA AT	16 319 15 320
p82w93	CCT GTC AAC ATA ATT A CCT GTC AAC ATA ATT	16 321
p82w94 p82w95	CCT GTC AAC ATA ATT G	16 322
P82w96	CCT ACA CCT GTC AA	14 323
p82w97	T GTC AAC ATA ATT GG	15 324
p82w98	T GTC AAC ATA ATT GGA	16 325
£		

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## Table 3 - Cont'd-9

p82m99 p82m100 p82m101 P82m102		Т	ACA	CCT CCT	TTC ATC	AAC	ATA ATA ATA	ATG			16 16 17 16	326 327 328 329
p82m103				CCT	GCC		ATA				16	330
p82m104			ACA	CCT	GCC		ATA	AG			16	331
p82m105				CCC		AAC	ATA	7.0			15 15	332 333
p82m106		-	CG	CCC	TTC		ATA	AG			15 15	333
p82m107		T	ACG ACA	CCC	GTC		AT				14	335
p82w108		CT CCT	ACA		GTC	AAC					14	336
p82w109 p82w110		CCI				AAC	АТА	ATG			15	337
p82w110						AAC					16	338
p82w112		CT	ACA		GTC	AAC					14	339
p82w113		CT		CCA	GTC	AAC	A				15	340
p82w114			ACA		GTC	AAC					15	341
p82w115			ACA			AAC		AG			16	342
p82w116		T	ACG	CCT	GTC						15	343
p82w117				CCT		AAC					15	344
p82w118		T	ACG	CCT	GTC	AAC	A				14 15	345 346
p82m119		CCT		CCT	TTC TTC	AAC AAC					14	347
p82m120	Α	CT CCT		CCT	TTC	AAC					15	348
p82m121 p82w122	A	CCI	ACG	CCT	GTC		ATA	AGG			16	349
p82w122		Т	ACG	CCT	GTC	AAC	ATA				16	350
p82w124		_	CG	CCT	GTC		ATA	AGG			15	351
p82m125		Τ	ACA	CCT	TTC	AAC	GTA				16	352
p82m126			ACA	CCT	TTC	AAC	GTA	AGG			16	353
p82m127			CA			AAC		ATG			16	354
p82m128			A	CCT	TTC			ATT	~~~	7.07	16	355
p82o129					C	AAC		ATT		AGA	16 <b>1</b> 5	356 357
p82o130					C	AAC	GTA	ATT	GGA	AG	1.0	337

Table 3 - Cont'd-10

	86	87	88	89	90	91	92	93	94	length	Seq ID
D00 1	GGA	AGA					CAG	TTA	GGT	1.0	250
P90w1			AAT			ACT ACT				16 17	358 350
P90w2			AAT AAT				CAG	7.00		18	359 360
P90w3		_	AAT	CTG			CAG			17	361
P90w4 P90w5		AGA		CTG						19	362
P90w5				CTG						20	363
P90w7				CTG						21	364
P90w8	Δ.	SA AF								20	365
P90w9		AGA								21	366
P90w10		AGA								21	367
P90m11		AGA					CAG			20	368
P90m12		AGA		CTG						21	369
P90m13	A	AGA								20	370
P90m14		AGA		CTG						21	371
P90m15	Α	AGA	AAT	CTG	ATG	ACT	CAG	ATG		21	372
P90m16	GA	AGA	AAT				CAG	ATT		20	373
P90m17	GGA	AGA	TAA			ACT				21	374
P90m18	A	AGA				ACT				19	375
P90m19				CTG						21	376
P90m20			AAT	CTG					G	20	377
P90m21			AAT				CAG		G	20	378
P90m22		A	AAT	CTG						19	379
P90m23		70.	TAA		ATG		CAG			18	380
P90w24			AAT		TTG		CAG		G	20	381
P90w25		A	AAT AAT	CTG			CAG		<u></u>	19 19	382 383
P90w26 P90w27			AAT				CAG	CII	G	14	384
P90w27			AAT		TTG		CAG			15	385
P90w29		Δ	AAT			ACT				15	386
P90w30						ACT				16	387
P90m31		7.7	AAT			ACT				14	388
P90m32			AAT			ACT				15	389
P90m33		А	AAT			ACT				15	390
P90m34		A	AAT	CTG	ATG	ACT	CAG			16	391
P90w35		GA	AAT	CTG	TTG	ACT	С			15	392
P90w36		GΑ	ACT	CTG			С			15	393
P90w37			${ m T}$				CAG	ATG		15	394
P90w38			AAT			ACT				15	395
P90w39				CTG						15	396
P90w40		A		CTG						15	397
P90w41				CTG						15	398
P90m42		ית		CTG						15 1 =	399
P90m43		A	TAA TA				CA	7) (		15 15	400
P90w44 P90w45			AT				CAG CAG			15 15	401 402
P90w45		A C A	חים ע	CTG			CAG	WII		15	402
P90m47		AUA		CTG			CAG	AG		15	404
					0						- 0 1

Table 3 - Cont'd-11

P90m48 P90m49	AGA AAT C		Т	15 17 15	405 406 407
P90w50	AAT A			16	408
P90w51	GA AAT A			15	409
P90w52	AAT T			16	410
P90w53	GA AAT T			15	411
P90w54	AAT A				412
P90w55	A AAT A'			15	
P90m56		G ATG ACC CAG		15	413
P90m57	A CAG A			15	414
P90w58	AAC A			15	415
P90w59		rg ttg act cag		15	416
P90w60		TG TTG ACT CAG CT		14	417
P90w61		TG TTG ACT CAG C		14	418
P90m62		CT ATG ACT CAG CT		14	419
P90m63	C	IG ATG ACT CAG C-		14	420
P90w64		IG ACT ACA CAG C	ΓT	14	421
P90w65	C	IG TTG ACA CAG C	-G	14	422
P90w66	AAT C	TG TTG ACA CAG		15	423
P90w67	AAC C	TG TTG ACT CA		13	424
P90w68	A AAC C	IG TTG ACT C		13	425
P90w69	GA AAC C	TG TTG ACT		13	426
p90w70		TG TTG ACT CAG A'	TT G	15	427
p90w71		TG TTG ACT CAG A		16	428
p90w72		G TTG ACT CAG A	TT GGG	15	429
p90w73		10 11	TT G	15	430
p90w74		TG TTG ACA CAG C		15	431
p90w75			TT GGG	15	432
p90w76		TG TTG ACT CAG C	TT G	15	433
p90w77			TG	15	434
p90w78		G TTG ACT CAG C	TT G	14	435
p90w79			TT G	15	436
p90w80		G TTG ACC CAG A	TT G	14	437
p90w81		G TTG ACC CAG A	TT GGG	15	438
p90m82		TG ATG ACT CAG A	TT G	15	439
p90m83		TG ATG ACT CAG A	TT GGG	16	440
p90m84		G ATG ACT CAG A	TT GGG	15	441
p90m85		G ATG ACT CAG A	TT GGT	16	442
p90m86		CTG ATG ACT CAG	CTT	15	443
p90m87		TG ATG ACT CAG	CTT G	15	444
P90w88	TAA A	CTG TTG ACT CA		15	445
P90w89	TAAA	CTG TTG ACT CA		15	446
p90w09	TAAA	CTG TTG ACT CA		15	447
p90w90 p90w100	דעע	CTG ATG ACT CAG		15	448
p90w100 p90m92	דעג ע	CTG ATG ACT CA		16	449
p90m93	GA AAT	CTG ATG ACT C		15	450
p90m94	011 1111	IG ATG ACT CAG A	<b>r</b> G	15	451
p90m95	AGA AAT			15	452
p90m96	A AGA AAT	ATG ATG ACT		16	453
Padilao					

Table 3 - Cont'd-12

p90m97 A p90m98 A p90m99 p90m100 p90m101 p90m102	AGA	AAT AAC	ATA ATA ATG CTG	ATG ATG ATG ATG ATG	ACT ACT ACC ACT ACT	CAG CAG C			16 16 16 15 15	454 455 456 457 458 459
p90m103	A		TTG CTG		ACT ACT	ATG	ACT		16 14	460 461
p90m104 p90m105		AC			ACT	CAG	A		16	462
p90m106		TA	CTG	ATG	ACT	CAG	ATG		16 14	463 464
p90m107		TA	CTG CTG	ATG ATG	ACT ACT	CAG CAG	ATT	G	16	465
p90m108 p90m109	AGA	TAA	CTG	ATG	ACT	C			16	466
p90m110	AGP	TAA	CTG		ACT				15 15	467 468
p90m111 G		TAA A	CTG	ATG ATG					16	469
p90m112 GGZ p90m113 GZ		AAT	CTG						16	470
p90m114		TAA A	CTG			C7\ C			14 15	471 472
p90w115		TAA T	CTG CTG		ACT	CAG CAG			16	473
p90w116 p90w117		TA			ACT				15	474
p90w118		TAA A							16 15	475 476
p90w119	G	TAA A TAA			ACT ACT	CAG	r T		15	477

Table 5

	non-B	86	7	٧	ዞ '	ဖ	0		
	Type B non-B	95.7	1 1	H C	0.0	т. Т.	1.1		
probes for	codon p50	w3.1	V V***	ተ ( ታ ፤ >	M52	m3.7	משנו		_
	non-B	7.0	> (	77	4	c	o 0	0	
	Type B	117 2	6.1	11.7	91	, ,	7.0	<b>&gt;</b>	
	probes tor	codon pro	w4./	w45	C772	7 7	m4 1	neg.	
-	ſ	ype B non-B	8	0	•	4	0	0	<del>,  </del>
		H	95.7	-	T	<b>⊣</b>	1.1	H.	_
•	probes for	codon p30	w2.5	1 0	W 2.3	w32	w36	m23	Š

Table 5 - Cont'd

17.5 0 65 2.5 2.5 22.5 10 5 2.5 10 5
Type B 50 66.1 7.1 12.5 7.1 5.4 5.4 0 0 19.6
probes for codon p90  w27  w37  w39  w50  w52  w69  w73  w73  m43  m56  neg.
100 10 12 0 10 0 0 0 0 0 0 0 2 0 0 12 0 0 10 0 10 10 10 10 10 10 10 10 10 10
Type B 81.9 2.1 1.1 1.1 4.3 2.1 1.1 1.1 1.1 1.1 3.2
probes for codon p82/84 w91 w60 w111 w89 w42 m36 m67 m38 m107 m40 m63 m101 neg.
non-B 48 62 18 0 8 4 4 4 4
Type B 71.3 81.9 3.2 6.4 4.3 0 0 3.2 14.9
probes for codon p54  w3  w34  w14  w19  w22  w26  w27  m35  m37  neg.

### Table 6

p30	USA	France	U.K.	Brazil	Spain	Luxemb.	Belgium
w25	98.9	99.4	88.9	98.3	94.3	100.0	97.0
<b>w</b> 29	2.5	0.6	0.0	1.7	0.0	0.0	0.0
w32	3.3	0.6	5.6	5.2	5.7	6.7	1.5
w36	2.5	0.0	0.0	3.4		0.0	1.0
m23	3.1		0.0	0.0		0.0	1.0
neg.		0.6	5.6	0.0	0.0		1.0
******							
p46/48	USA	France	U.K.	Brazil	Spain	Luxemb.	Belgium
w47	94.2		83.3		97.1		82.9
w45	8.6	15.6	0.0	1.7			11.1
w72	4.2		16.7	0.0	2.9		
m41	0.0		0.0				
neg.	2.8	4.5	0.0	0.0		0.0	2.5
2							
p50	USA	France	U.K.	Brazil	Spain	Luxemb.	Belgium
w31	96.4	97.4	100.0	96.6	100.0	100.0	96.5
w44	1.7	0.6	0.0	1.7	0.0	0.0	1.0
w52	10.0	4.5	0.0	1.7	2.9	6.7	9.0
m37	2.5	0.6	0.0	1.7	0.0	6.7	0.5
neg.	3.1	2.6	0.0	3.4	0.0	0.0	1.5
p54	USA	France	U.K.	Brazil	Spain	Luxemb.	Belgium
w34	96.9	82.5	97.2	87.9	_		89.4
w3	84.7	77.9	94.4	69.0	82.9	46.7	76.9
w14	3.3	5.8	0.0	3.4	11.4	0.0	6.5
w19	9.2	2.6	0.0	1.7	2.9	6.7	5.5
w22	2.8	10.4	0.0	0.0	5.7	0.0	2.5
w26	0.0	1.3	0.0	0.0	0.0	0.0	0.0
w27	0.0	1.9	0.0	0.0	0.0	0.0	0.5
m55	0.0	0.0	0.0	0.0	0.0	13.3	0.5
m35	1.1	0.0	2.8	6.9	0.0	46.7	3.0
m37	0.0	0.0			0.0	13.3	0.0
neg.	0.6	1.3	0.0	1.7	0.0	0.0	2.0
p82/84	USA	France	U.K.	Brazil	Spain	Luxemb.	Belgium
w91	91.6	93.5	94.4	77.6	100.0	73.3	85.9
w60	6.4	2.6	0.0	1.7	2.9	13.3	5.5
w111	3.6	0.6	0.0			0.0	0.5
w89	7.0	1.9	0.0			0.0	3.0
w42	0.6	0.0	2.8			0.0	2.0
m36	0.3	0.0	0.0			0.0	0.0
m67	0.0	0.0	0.0	0.0	0.0	0.0	0.5

## Table 6 - Cont'd

m38 m105 m127 m40 m63 m101 neg.	0.0 0.0 0.0 2.8 0.3 1.9 2.5	0.0 0.0 0.0 0.0 0.0 4.5 3.9	0.0 0.0 0.0 8.3 0.0 0.0	0.0 0.0 0.0 3.4 1.7 3.4	0.0 0.0 0.0 5.7 2.9 0.0	6.7 0.0 0.0 46.7 13.3 6.7 6.7	0.0 0.0 0.0 0.5 4.0 5.0
p90 w27 w37 w39 w50 w52 w69 w73 w79 m43 m56	USA 51.1 91.9 0.0 2.6 8.4 5.2 6.1 7.1 1.9	France 45.5 73.4 0.0 23.8 11.2 1.4 9.1 11.2 0.0	U.K. 34.3 80.0 0.0 2.9 5.7 5.7 0.0 8.6 11.4 0.0	Brazil 47.7 81.8 0.0 13.6 6.8 2.3 0.0 9.1 0.0	Spain 52.8 88.9 0.0 11.1 13.9 0.0 8.3 5.6 0.0 0.0	Belgium 25.7 55.2 2.9 21.9 4.8 3.8 6.7 5.7 8.6 0.0	

	Table 7	Tm 1	engte	Seq ID
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pc30w25 pc30w29 pc30w32 pc30w36 pc30m23	26 27 28 29 30 31 32 33 34 35 ACA GGA GCA GAT GAT ACA GTA TTA GAA GAA GCA GAT GAT ACA GT A GCG GAT GAT ACA GCA GAT GAC ACA GCA GAC GAT ACA GT GCA GAC GAT ACA GT A GCA GAC GAT ACA GG A GCA GAT AAT ACA GT	40 36 42 40 40	14 13 14 14 15	31 35 38 42 29
pc48w37 pc48w47 pc48w73 pc48w45 pc48w72 pc48m41	44 45 46 47 48 49 50 51 52 CCA AAA ATG ATA GGG GGA ATT GGA GGT ATG ATA GGG GGA ATT AAA ATG ATA GGG GGA A AGA ATG ATA GGG G AAA ATG ATA GGG GGA A AAA ATG ATA GGG GGA A AAA ATG ATA GGG GGA ATG ATA GGG GGA ATG ATA GTG GGA ATT	42 42 42 40	15 15 14 18 16 15	512 93 513 91 120 87
pc50w31 pc50w44 pc50w52 pc50m37	48	42 42 40	15 14 14 12	151 164 172 157
pc54w34 pc54w14 pc54w19 pc54w22 pc54w26 pc54w27 pc54m35 pc54m37 pc54m55	51 52 53 54 55 56 57 58  GGA GGT TTT ATC AAA GTA AGA CAG  GA GGT TTT ATC AAA GT  A GGC TTT ATC AAA GTA  A GGC TTT ATC AAA GTA  GA GGT TTT ATT AAA GTA  A GGT TTC ATT AAA GTA  A GGT TTT ATT AAA GTA  GGT TTT GTC AAA GTA  GGT TTT GTC AAA GTA  A GGT TTT GTC AAA GTA  GGT TTT GTC AAA GTA  A GGT TTT GTC AAA GTA	42 42 42 42 42 40 40 42	16 16 16 17 16 16 15 15	212 189 194 197 202 204 213 215 516
pc82w91 pc82w60 pc82w111 pc82w89 pc82m101 pc82w42 pc82m38 pc82m10 pc82m10	ACA CCT GTT AAC ATA AG  ACA CCT ATC AAC ATA AT  CA CCT GTC AAC GTA  ACA CCT TTC AAC ATA  ACG CCC TTC AAC ATA		16 17 16 17 17 14 15 15	318 287 338 316 517 269 265 332 354

## Table 7 - Cont'd

pc82m40 pc82m63 pc82m36 pc82m67			CF ACF	roo <i>A</i> roo <i>A</i> roo <i>A</i>	GCC ACC	AA C	ATA	AG		44 42	15 16 15 14	267 290 518 519
	86	87	88	89	90	91	92	93	94			
	GGA	AGA	AAT	CTG	TTG	ACT	CAG	ATT	GGT			
pc90w27			AAT	CTG	TTG	ACT	CA			38	14	384
pc90w37			T	CTG	TTG	ACT	CAG	$\mathtt{AT}$			15	514
pc90w39		GΑ	GTC	AAC	AGA	GTT	С				15	515
pc90w50			AAT	ATG	TTG	ACT	CAG			40	15	407
pc90w52			AAT	TTG	TTG	ACT	CAG			40	15	409
pc90w69		GΑ	AAC	CTG	TTG	ACT				40	14	426
pc90w73				ΤG			CAG			44	15	430
pc90w79				ΤG	TTG	ACC	CAG	ATT	G	44	15	436
pc90m138		GT	CAT	C AG	TTT A	T CT					14	510
pc90m56			AAT	ATG	ATG	ACC	CAG			42	15	413

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#### **CLAIMS**

- 5 1. Method for determining the susceptibility to antiviral drugs of HIV viruses in a biological sample, with said method comprising:
  - a) if need be, releasing, isolating or concentrating the polynucleic acids present in the sample;
  - b) if need be amplifying the relevant part of the protease gene of HIV with at least one suitable primer pair;
  - c) hybridizing the polynucleic acids of step a) or b) with at least one of the following probes:

probes specifically hybridizing to a target sequence comprising codon 30; probes specifically hybridizing to a target sequence comprising codon 46 and/or 48; probes specifically hybridizing to a target sequence comprising codon 50; probes specifically hybridizing to a target sequence comprising codon 54; probes specifically hybridizing to a target sequence comprising codon 82 and/or 84; probes specifically hybridizing to a target sequence comprising codon 90; or the complement of said probes;

- further characterized in that said probes specifically hybridize to any of the target sequences presented in figure 1, or to the complement of said target sequences;
- d) inferring from the result of step c) whether or not a mutation giving rise to drug resistance is present in any of said target sequences.
- 25 2. Method according to claim 1, further characterized in that said polynucleic acids of step a) or b) hybridize with at least two of the said probes, or to the complement of said probes.
- 3. Method according to claim 2, further characterized in that said probes are chosen from the following list: seq id no 7 to seq id no 477, seq id no 510 to seq id no 519 or the complement of said probes.
  - 4. Method according to any of claims 1 to 3, further characterized in that said primer pair is chosen from the following primers: seq id no 3, seq id no 503, seq id no 504, seq id no 4, seq id no 506, seq id no 507, seq id no 508 and seq id no 509.
  - 5. Method according to any of claims 1 to 3, further characterized in that:

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step b) comprises amplifying a fragment of the protease gene with at least one 5'-primer specifically hybridizing to a target sequence located at nucleotide position 210 to 260 of the protease gene, in combination with at least one suitable 3'-primer, and step c) comprises hybridizing the polynucleic acids of step a) or b) with at least one of the probes specifically hybridizing to a target sequence or its complement, comprising codon 90.

- 6. Method according to any of claims 1 to 3, further characterized in that:
  step b) comprises amplifying a fragment of the protease gene with at least one 3'-primer specifically hybridizing to a target sequence located at nucleotide position 253 (codon 85) to position 300, in combination with at least one suitable 5'-primer, and step c) comprises hybridizing the polynucleic acids of step a) or b) with at least one of the probes specifically hybridizing to a target sequence or its complement, comprising any of codons 30, 46, 48, 50, 52, 54, 82 and 84.
- Method according to claim 5, further characterized in that the 5'-primer is seq id 5 and the 3'primer is one primer or a combination of primers chosen from the following primers: seq id no
  4, seq id no 506, seq id no 507, seq id no 508 and seq id no 509.
- 8. Method according to claim 6, further characterized in that the 5'-primer is one primer or a combination of primers chosen form the following primers: seq id no 3, seq id no 503, seq id no 504 and the 3'-primer is seq id no 6.
  - 9. A probe as defined in any of claims 1 to 3, for use in a method for determining the susceptibility to antiviral drugs of HIV viruses in a biological sample.
  - 10. A nucleic acid comprising a nucleotide sequence represented by any of the following SEQ ID numbers: SEQ ID NO 478, SEQ ID NO 479, SEQ ID NO 480, SEQ ID NO 481, SEQ ID NO 482, SEQ ID NO 483, SEQ ID NO 484, SEQ ID NO 485, SEQ ID NO 486, SEQ ID NO 487, SEQ ID NO 488, SEQ ID NO 489, SEQ ID NO 490, SEQ ID NO 491, SEQ ID NO 492, SEQ ID NO 493, SEQ ID NO 494, SEQ ID NO 495, SEQ ID NO 496, SEQ ID NO 497, SEQ ID NO 498, SEQ ID NO 499 and SEQ ID NO 500; or a fragment thereof, wherein said fragment consists of at least two contiguous nucleotides and contains at least one polymorphic nucleotide.
- A primer as defined in any of claims 4 to 8, for use in a method for determining the susceptibility to antiviral drugs of HIV viruses in a biological sample.

- 12. A diagnostic kit enabling a method for determining the susceptibility to antiviral drugs of HIV viruses in a biological sample, with said kit comprising:
  - a) when appropriate, a means for releasing, isolating or concentrating the polynucleic acids present in said sample;
  - b) when appropriate, at least one of the primers of any of claims 4 to 6;
  - c) at least one of the probes of any of claims 1 to 3, possibly fixed to a solid support;
  - d) a hybridization buffer, or components necessary for producing said buffer;
  - e) a wash solution, or components necessary for producing said solution;
  - f) when appropriate, a means for detecting the hybrids resulting from the preceding hybridization;
  - h) when appropriate, a means for attaching said probe to a solid support.

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Figure 1

### Codon 30

### Codon 46/48

### Codon 50

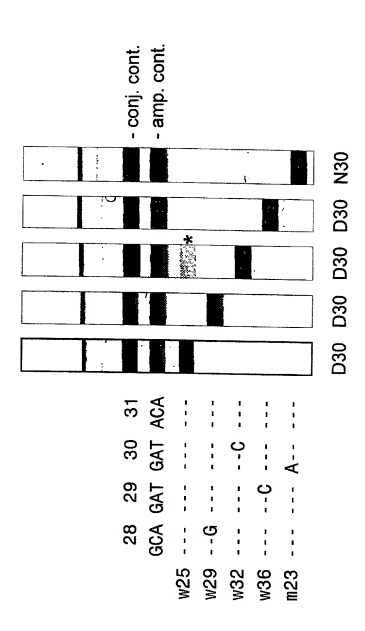
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2/21

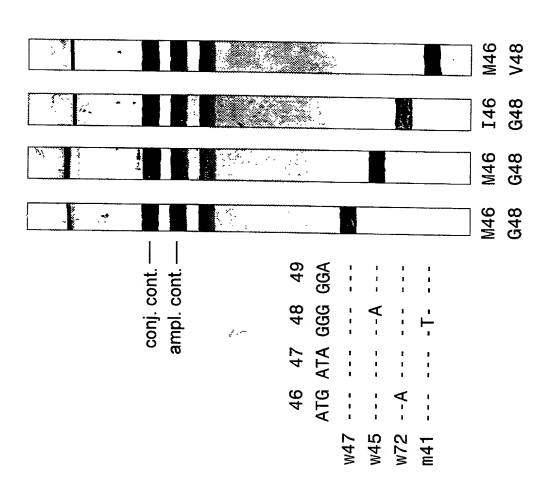
Figure 1 - Cont'd

### Codon 82/84

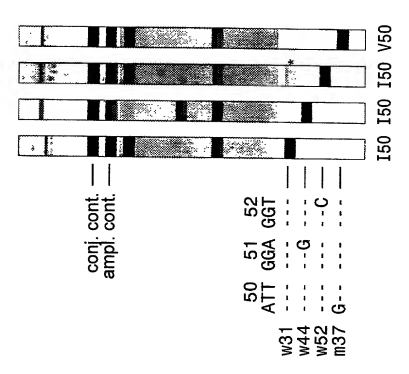
### Codon 90



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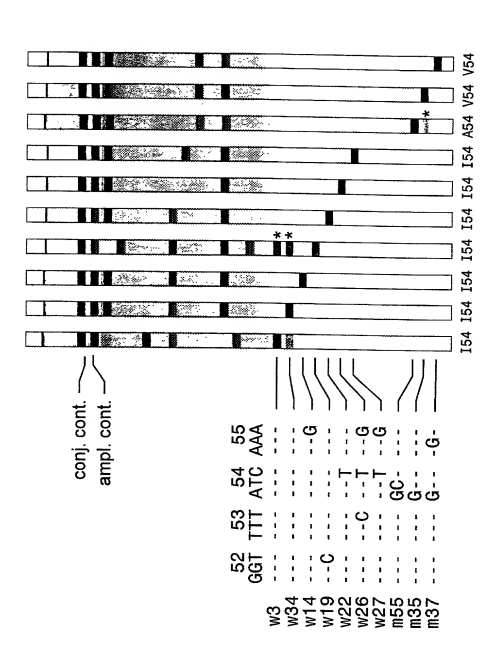
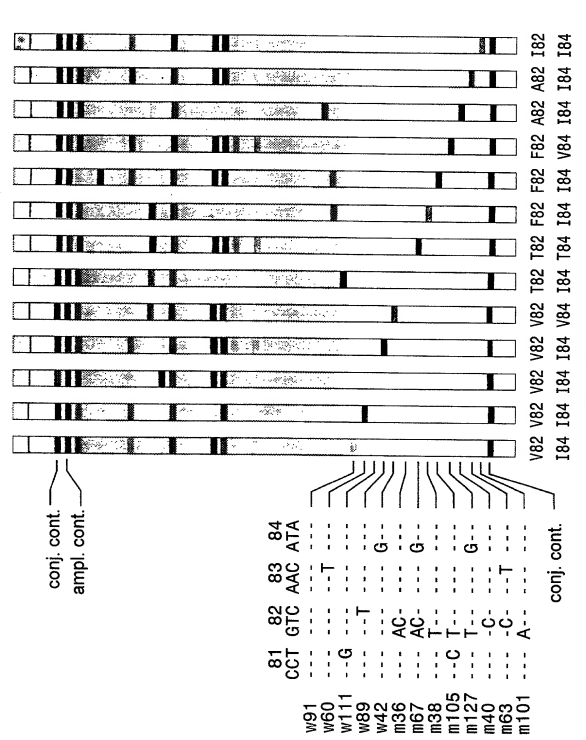
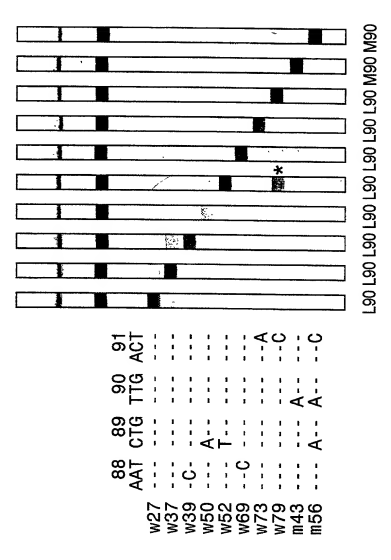


Figure 2D

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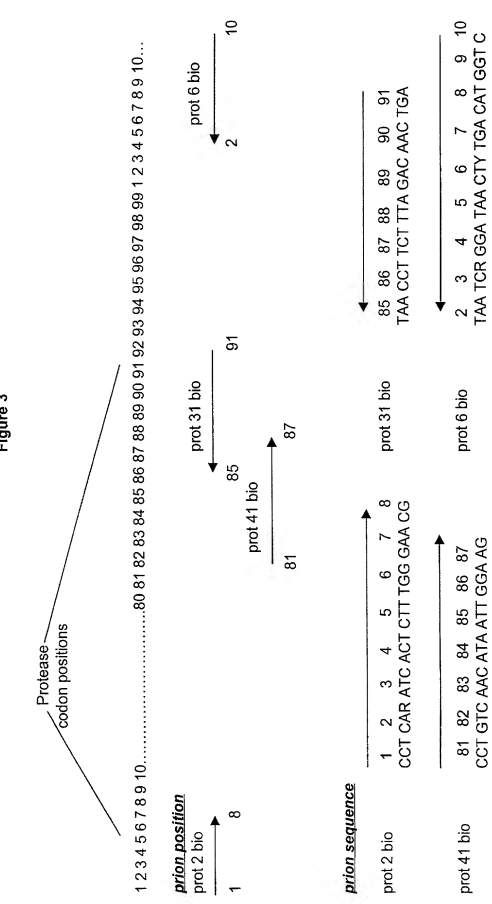


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igure 21

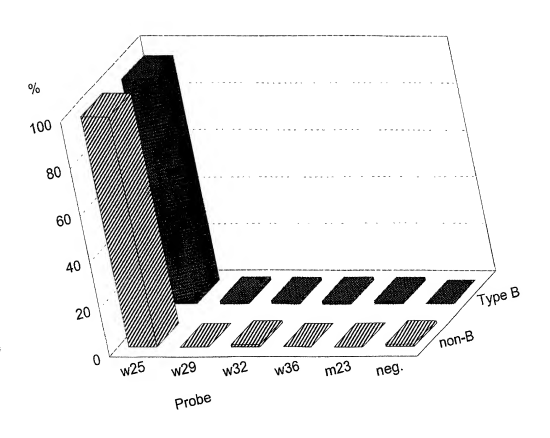




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Figure 4A



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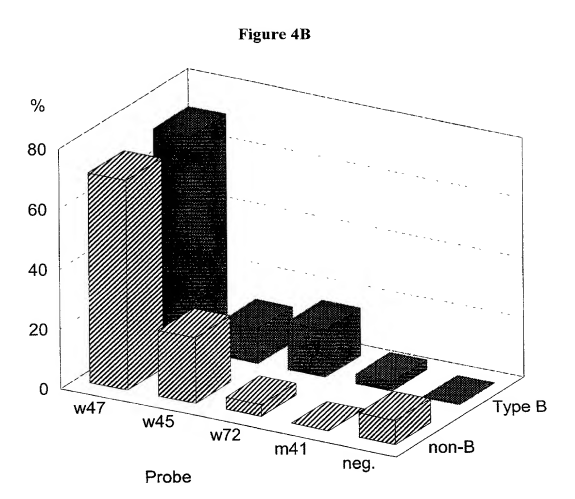
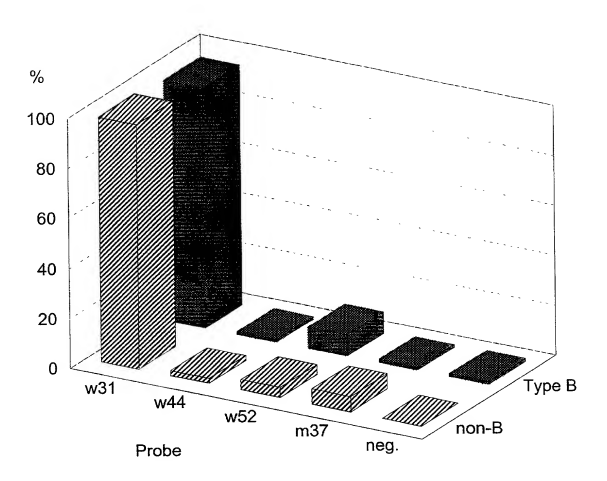
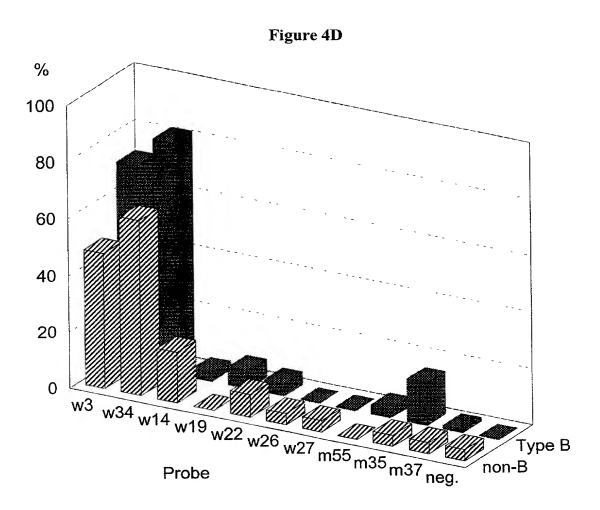


Figure 4C

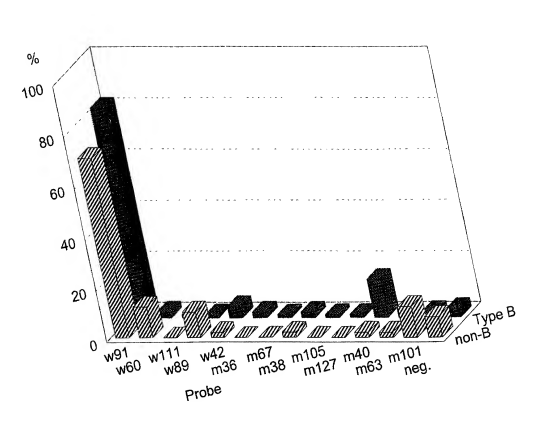




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Figure 4E



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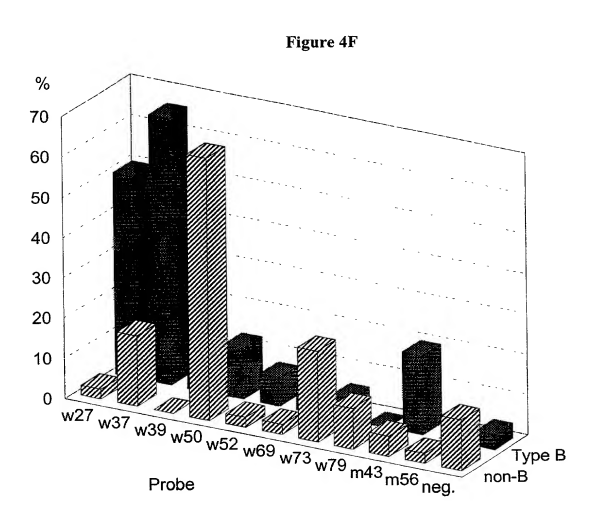
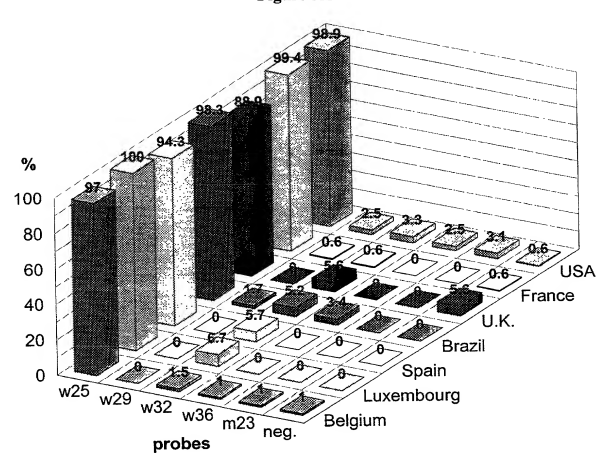
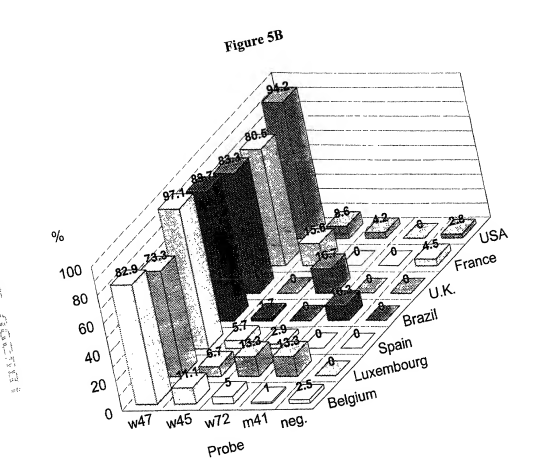


Figure 5A



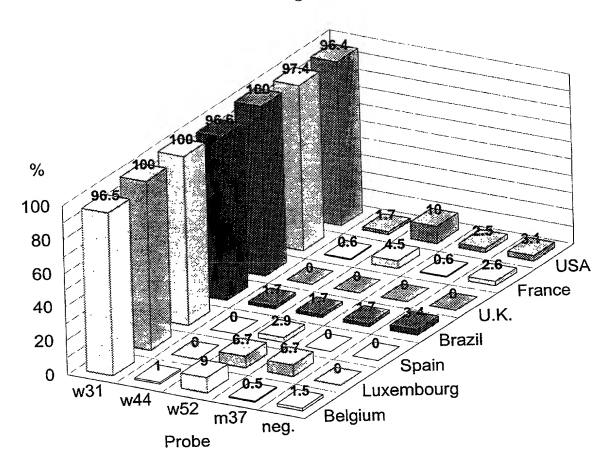
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Figure 5C

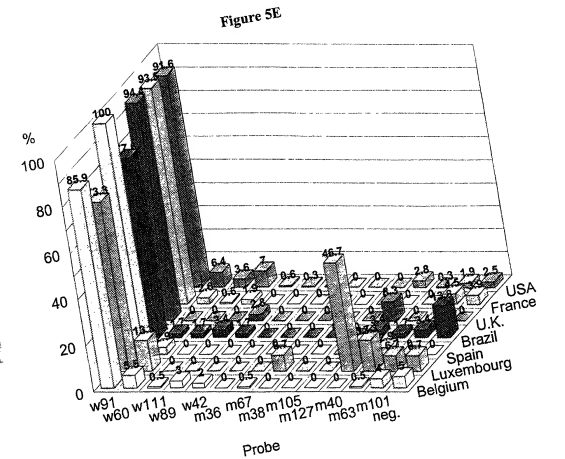


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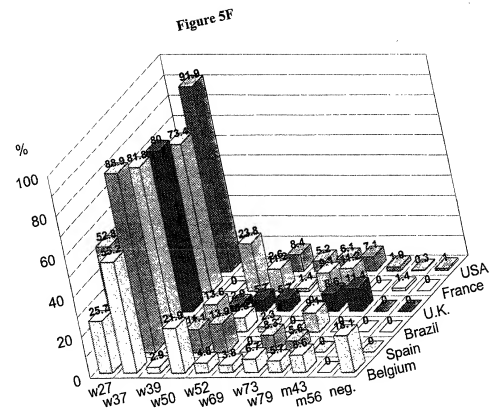




Probe

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Probe

## DECLARATION

As below named inventors, we hereby declare that:

Our residence, post office address and citizenship are as stated below next to our names.

The below named inventors are the original, first and joint inventors of the subject matter which is claimed and for which a patent is sought on the invention entitled **METHOD FOR DETECTION OF DRUG-SELECTED MUTATIONS IN THE HIV PROTEASE GENE**, the specification of which was filed as PCT International Application No. PCT/EP99/04317 on June 22, 1999 and accorded U.S. Serial Number \_\_\_\_\_\_\_.

We hereby state that we have reviewed and understand the contents of the above identified specification, including the claims.

We acknowledge the duty to disclose to the Patent and Trademark Office all information known to us to be material to patentability of the subject matter claimed in this application, as "materiality" is defined in Title 37, Code of Federal Regulations, § 1.56.

We hereby claim foreign priority benefits under Title 35, United States Code, § 119 (a)-(d) of any foreign application(s) for patent listed below and have also identified below any foreign application for patent or inventor's certificate having a filing date before that of the application on which priority is claimed.

## PRIOR FOREIGN APPLICATION(S)

**Priority Claimed** 

98870143.9	EP	24 June 1998	Yes
(Number)	(Country)	(Date Filed)	

We hereby claim the benefit under Title 35, United States Code, § 120 of any United States application(s), or § 365(c) of any PCT International application designating the United States of America, listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in the prior United States or PCT International application in the manner provided by the first paragraph of Title 35, United States Code, § 112, we acknowledge the duty to disclose all information known to me to be material to patentability of the subject matter claimed in this application, as "materiality" is defined in Title 37, Code of Federal Regulations, § 1.56, which become available between the filing date of the prior application and the national or PCT international filing date of this application.

PCT/EP99/04317	June 22, 1999	
(International Application No.)	(International Filing Date)	

We hereby direct that all correspondence and telephone calls be addressed to:

Patricia A. Kammerer
Howrey Simon Arnold & White, LLP
750 Bering Drive
Houston, TX 77057-2198
(713) 787-1438

Page 1 of 2

INNS:030 Declaration of Lieven STUYVER

attorneys for the assignee of this application.

WE HEREBY DECLARE THAT ALL STATEMENTS MADE OF OUR OWN KNOWLEDGE ARE TRUE AND THAT ALL STATEMENTS MADE ON INFORMATION AND BELIEF ARE BELIEVED TO BE TRUE; AND FURTHER THAT THESE STATEMENTS WERE MADE WITH THE KNOWLEDGE THAT WILLFUL FALSE STATEMENTS AND THE LIKE SO MADE ARE PUNISHABLE BY FINE OR. IMPRISONMENT, OR BOTH, UNDER SECTION 1001 OF TITLE 18 OF THE UNITED STATES CODE AND THAT SUCH WILLFUL FALSE STATEMENTS MAY JEOPARDIZE THE VALIDITY OF THE APPLICATION OR ANY PATENT ISSUED THEREON.

Inventor's Full Name	Lieven (nmi) STUYVER		
Inventor's Signature			
Date: Jan 10,	2001	Country of Citizenship:	BELGIUM
Holestraat 8 B-9552 Herzele BELGIUM ZEX			
Post Office Address, same as above if different from above			

## SEQUENCE LISTING

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